



Numerical simulation of compressible two-phase flow and reactive transport in porous media - Applications to the study of CO₂ storage and natural gas reservoir.

Irina Sin

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Irina Sin. Numerical simulation of compressible two-phase flow and reactive transport in porous media - Applications to the study of CO₂ storage and natural gas reservoir.. Earth Sciences. Ecole Nationale Supérieure des Mines de Paris, 2015. English. NNT : 2015ENMP0058 . tel-01306860

HAL Id: tel-01306860

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École doctorale n°398 :
Géosciences, Ressources Naturelles et Environnement

Doctorat ParisTech

T H È S E

pour obtenir le grade de docteur délivré par

l'École nationale supérieure des mines de Paris

Spécialité « Hydrologie, hydrogéologie quantitatives »

présentée et soutenue publiquement par

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le 8 décembre 2015

Numerical simulation of compressible two-phase flow and reactive transport in porous media — Applications to the study of CO₂ storage and natural gas reservoirs

Modélisation numérique d'écoulement diphasique compressible et transport réactif en milieux poreux — Applications à l'étude de stockage de CO₂ et de réservoir de gaz naturel

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Remerciements

I am grateful to École des Mines and BRGM for the financial support that made this work possible. I would also like to thank the MoMaS, GRC and SIAM organizations for their conferences and workshops.

Je remercie mon directeur de thèse, Vincent Lagneau, pour chacune des rencontres, éclairantes, chacune des explications, patientes et passionnantes, pour l'esprit scientifique qu'il m'a transmis, pour la liberté confiée et le temps consacré. Merci de m'avoir fait poser les problèmes et chercher de nouvelles possibilités. Спасибо за его и мои пérégrinations, заботу и за то, что был рядом в важные моменты моей жизни.

Je tiens à remercier Jérôme Corvisier d'avoir été mon maître de thèse. Je lui suis reconnaissante pour son soutien et pour m'avoir fait découvrir des visions alternatives sur des problèmes critiques.

I thank Denis Voskov for reading my thesis and providing valuable recommendations. I am grateful for his perceptive questions and fruitful discussions over the last two years. Я глубоко признательна Денису за советы, знания и за душевное общение.

I express my profound gratitude to Ulrich Mayer for reviewing the thesis, for his outstanding editorial contribution to this work, insightful questions and excellent discussion during the oral defense that disclosed new interesting issues to me.

Je tiens à remercier Roland Masson d'avoir accepté de présider mon jury de thèse. Je lui suis très reconnaissante pour toutes les connaissances qu'il m'a apportées depuis notre toute première rencontre, ce qui a joué un rôle important dans mes recherches.

Je tiens à remercier Mohamed Azaroual et Pascal Audigane d'avoir été mes encadrants. Je leur suis fortement reconnaissante pour l'accueil chaleureux au BRGM, pour les conseils, les échanges et l'expertise qui ont favorisé l'avancement de ce travail.

Je remercie vivement Christophe Coquelet pour la rédaction de la partie thermodyna-

mique ainsi que pour toutes les discussions productives, sa disponibilité et participation enthousiaste au travail, et pour m'avoir ouvert les portes du Centre Thermodynamique des Procédés de l'École des Mines (CTP). J'ai beaucoup apprécié bien la collaboration avec Martha Hajiw et Tianyuan Wang sur les données de CPA-PR. Mes remerciements vont également à Philippe Rivière du Centre Efficacité Énergétique des Systèmes (CES) pour ses conseils et son soutien permanent depuis le premier jour à l'École des Mines. Je remercie Claude Tadonki d'avoir partagé son expérience sur GMRES.

Je suis très reconnaissante aux murs et aux pavés de l'École des Mines et en particulier, à tout le Centre de Géosciences à Fontainebleau pour leur accueil cordial. J'ai beaucoup apprécié le travail sur le benchmark avec Laurent de Windt, son initiative et son expérience remarquable en géochimie. Je remercie Raphaël Mathieu d'avoir été mon guide dans le monde d'HYTEC, je le remercie également pour tout le support informatique, toujours immédiat. Mes remerciements vont bien entendu aussi à ma chère Rosa-Lynda, inestimable et indispensable, pour ses efforts administratifs et pour sa générosité d'âme.

Je tiens à remercier Anthony Michel pour ses conseils prudents et explications enrichissantes dès le début de ma thèse. J'ai tiré beaucoup d'enseignements de nos échanges sur le couplage et le benchmarking avec lui et Thibault Faney.

I am deeply grateful to Yalchin Efendiev for his precious advices and thoughtful discussions. I express my sincerest appreciation to Mikhail Panfilov, Alain Bourgeat, Leonid Pankratov, Sergey Shmarev, Tatyana Chechkina and Gregory Chechkin for the inspiration and delightful time we shared.

Je remercie également Caroline de Dieuleveult, Ghassan Jomaa, Anvar Farkhutdinov, Trevor Cox, Marco Campestrini, Tri Dat Ngo et Alain Gaunand pour les échanges et leur amitié.

Я от всего сердца признательна Оливье за его неустанную поддержку и доброту.

Мою дорогую и любимую Лену я благодарю за то, что была соучастницей в этом приключении.

Я благодарна моим родителям и сестре Дине за непоколебимую веру в меня, за все разговоры о диссертации, советы, заботу и любовь на протяжении всего пути, за присутствие сквозь время и расстояния.

할머니, 대륙과 섬에서 계시는 사랑하는 가족들 하늘처럼 끝없는 사랑과 격려를 주셔서 감사합니다 여러분 덕분에 제가 성장할 수 있는 것 같습니다 감사합니다.

Résumé

Les activités humaines dans la subsurface se développent rapidement (stockage de déchets, nouvelles techniques minières, stockage à haute fréquence de l'énergie), alors que les attentes du public et des autorités s'intensifient. L'évaluation de chaque étape de ces opérations souterraines nécessite des études détaillées de la sûreté et des impacts environnementaux. La modélisation multiphysique permet de comprendre et de prévoir le comportement des systèmes complexes, à différentes échelles de temps et d'espace.

Le but de ce travail est d'intégrer la résolution de l'écoulement diphasique compressible dans le cadre de codes de transport réactif à l'aide d'une méthode de séparation d'opérateurs. Un module multiphasique a été créé dans le code de transport réactif HYTEC. Une nouvelle approche a ensuite été développée pour coupler écoulement multicomposant multiphasique compressible, description de propriétés thermodynamiques des fluides, avec des codes de transport réactif. Une méthode alternative a été proposée pour résoudre séparément le transport multiphasique en utilisant des termes de taux de réaction numériques. Le système couplé a été vérifié et comparé à d'autres formulations de couplage pour la précision et la performance numérique. Un exercice de benchmark basé sur la modélisation d'un réservoir de gaz naturel a été proposé pour examiner les méthodes de couplage de simulateurs multiphasiques réactifs, leurs capacités numériques et caractéristiques.

Une partie de ce travail s'est focalisée sur la modélisation numérique et analytique de captage et stockage du carbone, l'impact de mélange convectif et de présence des impuretés dans le gaz injecté. La méthode de couplage a permis d'étudier des scénarios d'injection de gaz complexes, incluant sans s'y limiter des mélanges de quatre gaz, et de révéler de nouveaux comportements de la dynamique des fluides, gazeux et liquides — interaction entre le flux convectif densitaire et la séparation chromatographique.

Abstract

Human activity in the subsurface has been rapidly expanding and diversifying (waste disposal, new mining technologies, high-frequency storage of energy), while the public and regulatory expectations keep growing. The assessment of each step of underground operations requires careful safety and environmental impact evaluations. Multiphysics modeling provides an effective way to understand and predict the behavior of such complex systems at different time and space scales.

This work aims at incorporating a compressible multiphase flow into the conventional reactive transport framework by an operator splitting approach. A multiphase flow module was therefore implemented in the reactive transport software HYTEC using a new approach, developed to fully couple multiphase multicomponent compressible flow and description of the fluid thermodynamic properties with existing reactive transport codes. An alternative method based on the sequential iterative approach was then invented to separately model multiphase transport by means of numerical reaction rate terms. The coupled system was verified and compared with other coupling formulations for accuracy and computational performance. A benchmark exercise of modeling a natural gas reservoir was proposed to investigate the coupling methods of reactive multiphase simulators, as well as their numerical capacity and characteristics.

The final part of this work concerns the numerical and analytical modeling of carbon capture and storage and the impact of convective mixing and gas impurities in the injected stream in particular. The proposed coupling allowed to study complex gas injection scenarios, modeling a system including but not limited to four gases, and to reveal new behaviors of gas and liquid dynamics, specifically the interaction between density convective flux and chromatographic partitioning.

Résumé long

Les activités humaines dans la subsurface se développent rapidement (stockage de déchets, nouvelles techniques minières, stockage à haute fréquence de l'énergie), alors que dans le même temps les attentes du public et des autorités s'intensifient. L'évaluation de chaque étape de ces opérations souterraines nécessite des études détaillées de la sûreté et des impacts environnementaux. La modélisation multiphysique permet de comprendre et de prévoir le comportement des systèmes complexes, à différentes échelles de temps et d'espace. Il est important de développer une méthode efficace d'intégration de l'écoulement multiphasique dans le cadre de codes de transport réactif.

Dans ce travail, nous avons proposé une approche de couplage par séparation d'opérateurs qui a pour but de modéliser les systèmes chimiques multiphasiques en conservant la structure flexible d'un code de transport réactif existant et donc, elle retient le paradigme général de module de l'écoulement indépendant de réactions géochimiques. La méthode a été implémentée dans le code de transport réactif HYTEC ouvrant de nouvelles possibilités en termes d'applications et développements futurs.

Dans les codes de transport réactif, le nombre de composants dans le problème d'écoulement multiphasique est généralement minimisé puisque chaque composant supplémentaire augmente la dimension de système non-linéaire. Pour réduire la complexité numérique, la formulation de conservation des phases couplée au transport compositionnel linéaire et aux équations d'état cubiques s'est employée à résoudre l'écoulement multicomposant multiphasique compressible. Ainsi, plus le système chimique est riche en composants, plus le couplage d'écoulement peut être avantageux en comparaison avec les méthodes basées sur la formulation compositionnelle. La résolution d'écoulement peut être découplée comme un module à l'aide de la méthode proposée et intégrée dans le cadre de codes de transport réactif. Une approche alternative a ensuite été développée pour modéliser séparément le

transport multiphasique en utilisant des termes de taux de réaction numériques. Le système de couplage d'écoulement multiphasique et de transport réactif hérite la propriété de stabilité inconditionnelle du schéma complètement implicite d'écoulement, ce qui fait la modélisation des régimes d'advection dominante efficace.

Un problème de benchmark avec une solution auto-similaire a été modélisé pour vérifier la méthode. Par la suite, une comparaison avec une formulation globale implicite a été effectuée sur un problème de deux composants et deux phases, les résultats ont démontré la performance numérique de la méthode. Pour vérifier et valider les méthodes de couplage de simulateurs multiphasiques réactifs, le benchmarking est indispensable. Nous avons établi un exercice de benchmark basé sur la modélisation d'un réservoir de gaz naturel (avec trois gaz). Le problème suppose une relation étroite entre le transport et la géochimie et vise à examiner la capacité numérique des méthodes de couplage et leurs caractéristiques dans le contexte de l'écoulement et du transport multiphasiques multicomposants, solubilité de gaz et interaction de gaz-eau-roche.

Les équations d'état peuvent être employées globalement ou localement pour des propriétés thermodynamiques différentes au sein du couplage. La calibration de leurs paramètres varie en fonction des besoins. Un aperçu des équations d'état a été donné sous la forme d'une classification par type de modèle et son application à des problèmes différents. Une analyse comparative sur propriétés volumétriques de corps purs et de systèmes binaires a été effectuée.

Les développements numériques de ce travail et leur implémentation dans le code de transport réactif HYTEC ont été utilisés pour modéliser captage et stockage du carbone : en particulier, l'impact de mélange convectif et de présence des impuretés dans le gaz injecté. La méthode de couplage proposée a permis d'étudier des scénarios d'injection de gaz complexes, incluant sans s'y limiter des mélanges de quatre gaz, et de révéler de nouveaux comportements de la dynamique des fluides, gazeux et liquides — interaction entre le flux convectif densitaire et la séparation chromatographique, leurs effets sur la vitesse du front de courant de gaz et sur la dissolution globale de gaz.

Le problème de Hele-Shaw de mélange convectif dans la phase aqueuse en présence du gaz et de la frange capillaire a été modélisé. Les résultats ont indiqué qu'en plus des mouvements convectifs dans la phase aqueuse, les instabilités, qui continuaient à grandir sur la surface de contact irrégulière entre deux phases, ont engendré des perturbations cor-

respondantes dans la phase gazeuse, favorisant le mélange de gaz. Par la suite, le mélange convectif dans les deux phases a aussi été observé à l'échelle du réservoir.

La modélisation 3D de stockage du carbone montre une dissolution convective relativement faible à l'échelle du réservoir, puisque maillage à haute résolution est critique pour caractériser précisément le flux massique. Nous avons donc examiné la sous-estimation convective numérique sur la géométrie radiale en utilisant l'approximation par morceaux en temps d'une solution analytique, non-convective, couplée à une expression analytique de transfert convectif. La méthode donne une prédiction du rayon de courant en haut de l'aquifère en tenant compte de la dissolution convective. L'étude a confirmé que la modélisation 2D radiale pouvait être employée pour estimer le transfert convectif, en réduisant la dimension de maillage et en améliorant le mélange convectif en même temps.

Long abstract

Human activity in the subsurface has been rapidly expanding and diversifying (waste disposal, new mining technologies, high-frequency storage of energy), while the public and regulatory expectations keep growing. The assessment of each step of underground operations requires careful safety and environmental impact evaluations. Multiphysics modeling provides an effective way to understand and predict the behavior of such complex systems at different time and space scales. It is of great importance to develop an efficient method for integrating multiphase flow into the reactive transport framework.

In this work, we proposed an operator-splitting based coupling approach that is devised to model multiphase chemical systems, conserving the versatile structure of an existing reactive transport code, and hence, it retains the general paradigm of the flow module independent of the geochemical processes. The method was implemented in the reactive transport simulator HYTEC, building new capabilities in terms of applications and further developments.

In the reactive transport codes, the number of components in the multiphase flow problem is usually minimized since any additional component increases the dimension of nonlinear system. To reduce the computational complexity, the phase conservation formulation coupled with the linear component transport and cubic equations of state was employed for solving compressible multiphase multicomponent flow. Thus, the more the chemical system is abundant in components, the more the proposed flow coupling can be advantageous in comparison with the methods based on the compositional formulation. The flow solution can be split as a module, by means of the proposed method, and integrated into the reactive transport framework. Thereafter, an alternative way based on the sequential iterative approach was invented to separately model multiphase transport using numerical reaction rate terms. The coupled system of multiphase flow and reactive trans-

port inherits the unconditional stability of the fully implicit scheme of flow that allows to efficiently model advective dominant regimes.

The method was verified by modeling a benchmark problem admitting a self-similar solution. Then, a comparison with the global implicit formulation was carried out, that demonstrated the computational efficiency of the method on the two-phase two-component problem. In order to verify and validate coupling approaches of reactive multiphase simulators, benchmarking is essential. Thus, we presented a benchmark exercise with tightly connected transport and geochemistry by modeling a natural gas reservoir (with three gases) that aims at specifically analyzing the numerical capacity of the coupling methods and their characteristics in the context of multiphase multicomponent flow and transport, gas solubility and gas-water-rock interaction.

Within the coupling, the equations of state can be deployed globally or locally for different thermodynamic properties. Their parameters and formulations are calibrated according to the modeled property. An overview of equations of state was given as a classification by type of model and its applicability to different problems. A comparative analysis on volumetric properties of pure components and binary systems was carried out.

The numerical developments of this work and their implementation in the reactive code HYTEC, were then used to model carbon capture and storage and to study the impact of convective mixing and gas impurities in the injected stream. The proposed coupling allowed to investigate complex gas injection scenarios, modeling a system including but not limited to four gases, and to reveal new behaviors of gas and liquid dynamics: high interaction between density driven flow and chromatographic partitioning and their effects on the velocity of the front of gas current and on the overall gas dissolution.

The Hele-Shaw problem of convective mixing in the aqueous phase in the presence of the gas and the capillary fringe was modeled. It was shown that in addition to the convective motion in the aqueous phase, the instabilities growing on the rough contact surface between two phases implied corresponding perturbations in the gas phase, leading to the gas mixing. Afterward, the convective mixing in both phases was also observed at the reservoir scale.

The 3D modeling of carbon storage provides relatively lower convective dissolution at the reservoir scale, since a high-resolution grid is critical to accurately characterize the mass flux. We examined the numerical convective underestimation on the radial geometry

applying an analytical formulation that was obtained by time piecewise approximation of non-convective analytical solution coupled with an analytical expression of convective transfer. The method yields a prediction of current radius at the top of the aquifer with convective dissolution. The study exhibited that the 2D radial modeling could be efficiently used to estimate the convective transfer, reducing the grid dimension and enhancing the convective mixing at the same time.

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Chapter 1

Introduction

Les activités humaines dans la subsurface se développent rapidement, alors que dans le même temps les attentes du public et des autorités s'intensifient. L'évaluation de chaque étape de ces opérations souterraines nécessite des études détaillées de la sûreté et des impacts environnementaux qui reposent sur des simulateurs élaborés. Le développement de la modélisation multiphysique est donc nécessaire.

Différentes approches sont déjà mises en oeuvre pour réunir des fonctionnalités des simulateurs de réservoir et d'hydrogéochimie. Selon des principes similaires à la modélisation du transport réactif, il existe deux voies : l'approche globale implicite et l'approche par séparation d'opérateurs. La première est le choix de la majorité des simulateurs de réservoir qui sont, par construction, ciblés à résoudre de larges systèmes linéaires. Au contraire, l'extension du transport réactif à l'écoulement multiphasique est souvent effectuée par séparation d'opérateurs. Pour réduire la dimension du système, le problème d'écoulement ne consiste que de composants dominants ce qui limite son application. Il est donc important de développer une méthode efficace et généralisée d'intégration de l'écoulement multiphasique dans le cadre de codes de transport réactif qui permette de modéliser les systèmes multicomposants multiphasiques et les réactions chimiques complexes et de résoudre un système d'écoulement de dimension minimale. Dans ce travail, une approche par séparation d'opérateurs est proposée : la formulation de conservation des phases est couplée au transport compositionnel linéaire et aux équations d'état cubiques. En conséquence, la résolution de l'écoulement peut être découplée comme un module et intégrée dans des codes de transport réactif.

Le manuscrit commence par la description des formulations de l'écoulement multiphasique, le Chapitre 2. Après avoir choisi l'une d'elles, nous présentons l'implémentation d'un module d'écoulement diphasique, incompressible, immiscible, isotherme, dans le code de transport réactif HYTEC ainsi que des résultats numériques des modèles de référence à l'étape de vérification.

Le Chapitre 3 est consacré au couplage de l'écoulement multiphasique et du transport réactif. Tout d'abord, l'intérêt est porté aux méthodes de résolution de l'écoulement compositionnel, du transport réactif et de leur couplage. Par la suite, une nouvelle approche de la modélisation de l'écoulement multiphasique miscible compressible et son couplage avec le transport réactif sont développés et suivis de la comparaison des résultats numériques d'HYTEC et d'autres codes. Ainsi, un exercice de benchmark basé sur la modélisation d'un réservoir de gaz naturel est établi pour la vérification et validation des codes de couplage.

L'écoulement miscible compressible suppose la définition de propriétés thermodynamiques pour lesquelles les équations d'état peuvent être employées. Le Chapitre 4 montre les types de ces modèles. Puis nous avons mené une analyse comparative sur les propriétés volumétriques de corps purs et de systèmes binaires. Pour les propriétés des fluides, une liste succincte des modèles choisis est donnée.

Après l'implémentation des méthodes décrites et leur vérification, nous réalisons des simulations numériques de stockage du carbone, le Chapitre 5. L'étude commence par le problème 1D de la séparation chromatographique. Ensuite, le problème convectif avec une couche de gaz fine est traité. Nous continuons avec la modélisation 2D radiale du stockage de CO_2 et comparons les résultats numériques aux estimations analytiques. Le problème est étendu à l'injection de CO_2 avec des impuretés, trois scénarios d'injection sont modélisés.

Les conclusions et les propositions aux développements futurs sont présentées dans le Chapitre 6.

Human activity in the subsurface has been expanding and diversifying (waste disposal, new mining technologies, high-frequency storage of energy), meanwhile the public and regulatory expectations keep growing. Assessment of each step of underground operations including environmental impact evaluation, relies on elaborate simulators and gives rise to a strong necessity of developing multiphysics modeling. Reactive transport, a geochemical research and engineering tool, deals with multicomponent systems and complex chemical reactions. Extending it to the multiphase flow widens its applicability for understanding subsurface processes. We discuss below the issues of modeling reactive transport and multiphase flow in Sec. 1.1 followed by the outline of the thesis, Sec. 1.2. The symbols and abbreviations used in the document are detailed in Nomenclature, p. 187.

1.1 Modeling multiphase flow and reactive transport

Coupling the multiphase flow and reactive transport has already been performed both by the reservoir and hydrogeochemical modeling communities. The compositional flow simulators, based on implicit methods, are usually extended to the chemical modeling by solving the fully coupled system, since the framework is initially devised and aimed at large nonlinear systems: GEM-GHG (Nghiem et al., 2004), GPRS (Fan et al., 2012). By contrast, MoReS (Farajzadeh et al., 2012; Wei, 2012) and UTCHEM (Delshad et al., 2000) use an operator-splitting approach. The element-based version of GPRS (Fan et al., 2012) and MoReS, coupled with PHREEQC (Parkhurst et al., 1999), can solve the kinetic and equilibrium reactions in mineral phase. Only MoReS provides complex activity models and thermodynamic databases. Coupling existing codes is another alternative, it thus enforces the operator splitting routine; e.g., iCP (Nardi et al., 2014), the interface of PHREEQC and COMSOL is an ongoing project¹.

Several reactive transport simulators have succeeded in integrating multiphase flow by an operator-splitting approach: PFLOTRAN (Lu and Lichtner, 2007), STOMP (White and Oostrom, 2006), TOUGHREACT (Xu and Pruess, 1998). In these codes, the two-phase flow system is set for two components that are presented by water and air/CO₂/CH₄; the relevant thermodynamic/equation of state module should be chosen (A more detailed description of the coupling methods will be given in Sec. 3.1.3).

¹Á. Sáinz, personal communication, 2014.

In the reactive transport codes, the number of components in the multiphase flow system is usually minimized, in contrast to reservoir simulators. Adding a component increases the dimension of nonlinear multiphase flow system and therefore, the computational complexity of the problem. Let us suppose an isothermal multiphase flow problem of N_f fluid phases and N_c components. Hence, there are

- N_c nonlinear mass conservation equations for each component,
- $2N_f$ constitutive relations ($N_f - 1$ capillary pressure equations, 1 equation of the saturation sum, N_p equations of the mole/mass fraction sum),
- $N_c(N_f - 1)$ phase equilibria equations,

resulting in a problem of $2N_f + N_cN_f$ equations. Based on the Gibbs phase rule, the number of intensive variables is $N_c - N_f + 2$. Then, considering two phases, there are N_c unknowns, hence N_c nonlinear equations. Using a classic global approach (Coats, 1980), we solve N_c nonlinear mass conservation equations. The size of matrix arising from the linearization of this system, is $N_cN_{gr} \times N_cN_{gr}$, where N_{gr} is the grid dimension. Any additional component introduced to a multiphase flow problem indeed amplifies significantly computational complexity for high resolution modeling. Also, it requires the development of a thermodynamic module that is applicable to multicomponent systems and describes the fluid properties of real mixtures.

In order to avoid large flow systems, the formulation can be reduced by considering only dominant components, TOUGHREACT. The applications of this method are limited. A general method should be initiated to allow an arbitrary number of components, all the while preserving minimum dimension of the flow problem. This work proposes to employ a phase conservation formulation for solving compressible multiphase multicomponent flow by an operator-splitting approach. The phase formulation is coupled with the linear component conservation system, so that the flow module can be separated. Therefore, the dimension of multiphase system is independent of the number of components. The module can be integrated into the reactive transport framework as a conventional two-phase two-component or compositional flow module.

A new approach of multiphase modeling was implemented in the (SIA) reactive transport simulator, HYTEC (Lagneau and van der Lee, 2010a; van der Lee et al., 2003),

developed at Mines ParisTech. The biogeochemical reactions are performed by the speciation code CHESS (van der Lee, 2009). The hydrodynamic module of HYTEC accounts for saturated, unsaturated flow. The code was validated in different benchmarks (De Windt et al., 2003; Carrayrou et al., 2010; Lagneau and van der Lee, 2010b; Trotignon et al., 2005) and it was applied in numerous applications, such as cement degradation (De Windt L., 2010), radioactive waste disposal (Debure et al., 2013), geological storage of acid gases (Corvisier et al., 2013), uranium *in situ* recovery processes (Regnault et al., 2014).

1.2 Outline

Chapter 2 presents an overview of multiphase flow formulations. Then we describe the numerical implementation of one of them in HYTEC, supposing incompressible, immiscible, isothermal flow. The verification of the flow module is demonstrated on some reference models.

Chapter 3 is devoted to the coupling of the multiphase multicomponent flow and reactive transport. We give an overview on the numerical methods for each part of the problem and of the existing coupling approaches. Next, a new method of modeling compressible multiphase flow and its coupling with reactive transport are detailed. Also, an alternative way to separate multiphase transport modeling is given, by means of the reaction rate terms for the sequential iterative reactive transport framework. The first numerical results are then shown. Finally, we present an application (transformed into a benchmark proposition) that supposes a tight connection between transport and geochemistry by modeling a natural gas reservoir (with three gases), submitted to Mathematics and Computers in Simulation.

When modeling miscible flow, the choice of equation of state models can be decisive, especially at high pressure and temperature. In Chapter 4, the equation of state models are classified. For some of them, a comparative analysis on volumetric properties is carried out. The models of fluid properties are also discussed.

Chapter 5 concerns the numerical and analytical modeling of carbon capture and storage and the impact of gas impurity in the injected stream. The study starts from a 1D problem of chromatographic partitioning. Next, it is followed by a convective problem with a thin gas layer. Thereafter, we provide the numerical and analytical results of 2D

radial CO₂ modeling. The problem is then extended to three scenarios of CO₂ injection containing gas impurities, modeling a system with up to four gases.

Chapter 6 draws some conclusions and recommendations for future work.

Chapter 2

Multiphase Flow

Le Chapitre 2 introduit les notions de base de l'écoulement multiphasique, nécessaire pour comprendre la physique, établir le problème mathématique et ensuite le modéliser. Une fois les formulations mathématiques de l'écoulement données, nous pouvons choisir l'une d'elles et poser le problème de l'écoulement diphasique isotherme avec les hypothèses simplificatrices à cette étape — notamment les fluides sont supposés immiscibles, incompressibles. Les forces diffusives et les réactions chimiques sont négligées dans cette partie. Les méthodes numériques sont détaillées : les schémas de discrétisation en espace et en temps, la linéarisation et le jacobien analytique. La vérification est établie sur la modélisation d'un problème parabolique avec une solution auto-similaire et d'un problème hyperbolique de référence. Les résultats numériques sont comparés à ceux obtenus par d'autres codes et montrent également la convergence et la précision du module construit.

After introducing basic definitions related to hydrodynamics (Sec. 2.1), the general formulations of multiphase flow (Sec. 2.2) are presented in order to provide a foundation for the modeled processes and their further extension. One of these systems is then chosen for modeling under the hypotheses listed in Sec. 2.3. Next, the applied numerical methods are described in Sec. 2.4. The numerical results of water-flooding and of a self-similar solution are presented in Sec. 2.5. The diffusive flux and chemical reactions are not considered yet.

2.1 Definitions

Averaging over a representative elementary volume (REV), one can represent the fluid and rock properties passing from one region to another using a continuum approach. In this work, the macroscopic level is considered; i.e., approximately from 10^{-3} to 10 m. We first introduce notions of phase, porosity, saturation and component, then the Darcy-Muskat velocity is given with the description of fluid-matrix parameters of multiphase systems.

In case two or more fluids are immiscible (i.e., they are separated by physically distinct interfaces), each of the fluids is a phase and the system is multiphase. Fig. 2.1(a) illustrates two immiscible fluids in contact with a solid phase, where θ is the contact angle between the solid surface and the fluid-fluid interface curved under the surface tension. If $\theta < 90^\circ$, the fluid is called wetting, otherwise, non-wetting. Fig. 2.1(b) presents the interface in case of water/oil and gas/oil. If the two fluids are miscible (for instance, gases), there is no a sharp fluid-fluid interface, the fluids represent one phase. In this case, the phase consists of components: the system is multicomponent.

The porosity ϕ is the ratio of porous space within an REV relative to the total volume of the REV. The saturation S_α of phase α is the ratio of the volume occupied by the phase α in an REV relative to the total volume of pore space in the REV. Supposing that the fluids fill all porous space in the REV, the following relationship holds for the phase saturations:

$$\sum_{\alpha} S_{\alpha} = 1 \quad (2.1.1)$$

When fluid is displaced, a residual saturation $S_{\alpha r}$ is typically retained; i.e., the residual saturation of the wetting phase during drainage, the residual saturation of the non-wetting phase during imbibition. Thus, the effective saturation is defined as

$$\bar{S}_{\alpha} = \frac{S_{\alpha} - S_{\alpha r}}{1 - \sum_{\beta} S_{\beta r}}, \quad (2.1.2)$$

where α, β are the phases and then $\sum_{\alpha} \bar{S}_{\alpha} = 1$.

The composition of phase α can be described by mass/mole fractions, concentrations. The mass fraction $X_{\alpha,k}$ of the components k is the ratio of the mass of the component k to the total mass of the phase α , then

$$\sum_k X_{\alpha,k} = 1. \quad (2.1.3)$$

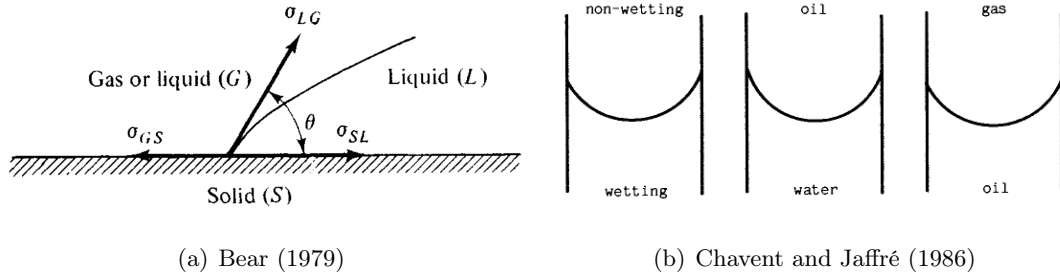


Figure 2.1 – Determination of the wetting and non-wetting phases.

By extension of Darcy's law, the velocity for multiphase flow Darcy-Muskat (Muskat et al., 1937) is expressed as follows

$$\mathbf{u}_\alpha = -\frac{k_{r\alpha}}{\mu_\alpha} \mathbb{K} (\nabla p_\alpha - \rho_\alpha \mathbf{g}), \quad (2.1.4)$$

where \mathbb{K} is the intrinsic permeability tensor, $k_{r\alpha}$ is the relative permeability of phase α , p_α is the pressure of phase α , ρ_α is the density of phase α , μ_α is the viscosity of phase α and \mathbf{g} is the gravity vector.

Due to the interfacial tension, the pressure changes from one phase to another and the difference is expressed by capillary pressure p_c :

$$p_n = p_w + p_c, \quad (2.1.5)$$

where p_n is the pressure of the non-wetting phase and p_w is the pressure of the wetting phase.

Since the porous space is filled by several fluids, the relative permeability of each phase $k_{r\alpha}$ represents a fraction of intrinsic permeability for phase α . The phase mobilities $\lambda_\alpha = \frac{k_{r\alpha}}{\mu_\alpha}$ are also used, the total mobility λ_{tot} is defined by

$$\lambda_{tot} = \sum_\alpha \lambda_\alpha. \quad (2.1.6)$$

The most common empirical functions of relative permeability $k_{r\alpha}$ and capillary pressure p_c are Brooks-Corey (1954) and van Genuchten (1980) models.

The Brooks-Corey capillary pressure function (Brooks and Corey, 1964) is given by

$$p_c(S_w) = p_b \bar{S}_w^{-\frac{1}{\lambda}}, \quad (2.1.7)$$

where p_b is the entry pressure, indicating a minimum non-wetting pressure required to enter into porous media saturated by the wetting phase; λ reflects pore size distribution, $[0.2, 3]$.

The van Genuchten capillary pressure model (van Genuchten, 1980) yields

$$p_c(S_w) = \frac{1}{\alpha} (\bar{S}_w^{-\frac{1}{m}} - 1)^{1-m}, \quad (2.1.8)$$

where $m = 1 - \frac{1}{n}$, $n > 1$.

The Brooks-Corey relative permeability model (Brooks and Corey, 1964) uses the function of capillary pressure (2.1.7) with the same λ :

$$k_{rw}(S_w) = \bar{S}_w^{\frac{2+3\lambda}{\lambda}}, \quad (2.1.9)$$

$$k_{rn}(S_n) = \bar{S}_n^2 (1 - (1 - \bar{S}_n)^{\frac{2+\lambda}{\lambda}}). \quad (2.1.10)$$

The van Genuchten relative permeability (van Genuchten, 1980) is expressed by

$$k_{rw}(S_w) = \bar{S}_w^\varepsilon (1 - (1 - \bar{S}_w^{\frac{n}{n-1}})^{\frac{n-1}{n}})^2, \quad (2.1.11)$$

$$k_{rn}(S_n) = \bar{S}_n^\gamma (1 - (1 - \bar{S}_n)^{\frac{n}{n-1}})^{\frac{2(n-1)}{n}}, \quad (2.1.12)$$

where $\varepsilon = 1/2$, $\gamma = 1/3$ (Helmig, 1997), $m = 1 - \frac{1}{n}$, $n > 1$, Mualem (1976).

2.2 Overview of multiphase flow formulations

We introduce briefly a survey of formulations for multiphase flow, that takes place according to the modeled problems and fluid properties. If the fluids are supposed to be immiscible, the general phase problem is solved, otherwise the compositional form should be applied to permit phase transition such as vaporization and dissolution. Detailed derivations of multiphase flow formulations are presented in Aziz and Settari (1979); Barenblatt et al. (1972); Bastian (1999); Bear (1972); Chavent and Jaffré (1986); Hassanizadeh and Gray (1979) and Peaceman (1977).

2.2.1 Immiscible multiphase flow

By considering the immiscible multiphase flow (no mass phase transfer), the mass conservation equation is written by analogy to single-phase flow but for each phase α , stating

the set of N_f partial differential equations

$$\frac{\partial(\phi S_\alpha \rho_\alpha)}{\partial t} + \nabla \cdot (\rho_\alpha \mathbf{u}_\alpha) = q_\alpha, \quad (2.2.1)$$

where q_α is the source term. Including the sum of saturations (2.1.1) and $N_f - 1$ capillary pressure equations (2.1.5), it constitutes a closed system of $2N_f$ equations for $2N_f$ unknowns. There exists several representations of system (2.2.1), so let us propose a concise list starting from the strongly coupled formulation to the weaker ones.

- Coupled general phase conservation equations (PS)

Substituting the velocity (2.1.4) in (2.2.1) results in the general phase formulation

$$\frac{\partial(\phi S_\alpha \rho_\alpha)}{\partial t} - \nabla \cdot (\rho_\alpha \lambda_\alpha \mathbb{K} (\nabla p_\alpha - \rho_\alpha \mathbf{g})) = q_\alpha. \quad (2.2.2)$$

For each phase α , a so-called natural variable, S_α or p_α , is chosen, leading to N_f unknowns. It is clear that these N_f nonlinear equations (2.2.2) are strongly coupled. Moreover, the nonlinear functions $k_{r\alpha}$ and p_c depend on the saturation. Supposing that p_w and S_n are the primary variables for the two-phase problem, the non-wetting phase conservation equation contains a term $\lambda_n \nabla p_c$. Choosing either the Brooks-Corey models (*BC*) for relative permeabilities (2.1.9), (2.1.10) and capillary pressure (2.1.7) or the van Genuchten models (*VG*) (2.1.11), (2.1.12) and (2.1.8) leads to the following singularities of $\lambda_n \nabla p_c$ (μ_α is neglected):

$$\begin{array}{ccc} & BC & VG \\ \lim_{S_n \rightarrow 0} k_{rn} p'_c(S_n) & = & 0 \quad 0 \\ \lim_{S_n \rightarrow 1} k_{rn} p'_c(S_n) & = & \infty \quad \infty \end{array} \quad (2.2.3)$$

Assuming (p_n, S_w) as the primary variables yields the appearance of $\lambda_w \nabla p_c$ in the wetting phase conservation equation and implies

$$\begin{array}{ccc} & BC & VG \\ \lim_{S_w \rightarrow 0} k_{rw} p'_c(S_w) & = & 0 \quad 0 \\ \lim_{S_w \rightarrow 1} k_{rw} p'_c(S_w) & = & < \infty \quad \infty \end{array} \quad (2.2.4)$$

Hence, any choice of one of these pairs entails restrictions at the extreme values of saturation in the presence of capillary forces. To avoid this problem, the pressure-pressure formulation can be used by extending the capillary pressure. In this case,

the phase disappearance problem arises that will be discussed later, p. 43. Numerical difficulties at the extreme values of saturation also come from the Jacobian structure, p. 21. As advantage of PS, the initial and Dirichlet boundary conditions have direct physical meanings.

The system (2.2.2) on (p_α, S_α) consists of one parabolic equation on the pressure (that turns to elliptic type in the incompressible case) and one degenerate parabolic equation on the saturation. If the capillary forces are neglected, the latter becomes hyperbolic, called Buckley-Leverett equation, and admits discontinuous solutions – shock waves. Since for the hyperbolic problems, several weak solutions can exist, the numerical scheme should satisfy an entropy condition (LeVeque, 2002). In Sec. 2.5.2, we demonstrate numerically that the Godunov method provides entropy-satisfying Riemann solutions.

- Decoupled pressure and saturation equations (DPS)

Decoupling eliminates the temporal derivative of saturation and leads to the pressure equation by summing the equations of system (2.2.2) divided by ρ_α

$$\frac{\partial \phi}{\partial t} + \sum_{\alpha} \frac{1}{\rho_{\alpha}} \left(\phi S_{\alpha} \frac{\partial \rho_{\alpha}}{\partial t} + \nabla \rho_{\alpha} \cdot \mathbf{u}_{\alpha} \right) + \nabla \cdot \mathbf{u} = \sum_{\alpha} q_{\alpha}. \quad (2.2.5)$$

Once the unknown for the pressure equation is chosen, the total velocity $\mathbf{u} = \sum_{\alpha} \mathbf{u}_{\alpha}$ and the saturation equation(s) can be reformulated. The phase pressure and global pressure formulations are discussed below.

- Phase pressure and saturation (DPPS)

Supposing the wetting phase pressure as primary variable allows to rewrite the velocities as

$$\mathbf{u} = -\lambda_{tot} \mathbb{K} (\nabla p_w + f_n \nabla p_c - \sum_{\alpha} f_{\alpha} \rho_{\alpha} \mathbf{g}), \quad (2.2.6)$$

$$\mathbf{u}_w = f_w \mathbf{u} + f_w \lambda_n \mathbb{K} (\nabla p_c + (\rho_w - \rho_n) \mathbf{g}), \quad (2.2.7)$$

$$\mathbf{u}_n = f_n \mathbf{u} - f_n \lambda_w \mathbb{K} (\nabla p_c - (\rho_n - \rho_w) \mathbf{g}), \quad (2.2.8)$$

where $f_{\alpha} = \lambda_{\alpha} / \sum_{\beta} \lambda_{\beta}$ is the fractional flow. Substituting the non-wetting phase velocity, for example, yields the saturation equation under the form of

mass conservation

$$\frac{\partial(\phi S_n \rho_n)}{\partial t} + \nabla \cdot (\rho_n \mathbf{u}_n) = q_n. \quad (2.2.9)$$

– Global pressure and saturation (GPS)

By defining the global pressure p such that $\nabla p = \sum_{\alpha} f_{\alpha} \nabla p_{\alpha}$, the total velocity simply becomes

$$\mathbf{u} = -\lambda_{tot} \mathbb{K}(\nabla p - \sum_{\alpha} f_{\alpha} \rho_{\alpha} \mathbf{g}). \quad (2.2.10)$$

Using the total velocity, the wetting phase saturation equation can be expressed by following

$$\frac{\partial(\phi S_w \rho_w)}{\partial t} + \nabla \cdot (\rho_w (f_w \mathbf{u} + f_w \lambda_n \mathbb{K}(\nabla p_c + (\rho_w - \rho_n) \mathbf{g}))) = q_w. \quad (2.2.11)$$

The main difference between the two formulations is the capillary pressure term $\lambda_n \nabla p_c$ that appears in the pressure equation of the DPPS formulation through the total velocity (2.2.6) as well as in the PS formulation (2.2.2). In the incompressible case, this nonlinearity is eliminated (2.2.10) by definition of the global pressure. When the compressibility is taken into account, the pressure equation (2.2.5) involves the phase velocities and then $\lambda_w \lambda_n \nabla p_c$, where $\lambda_w \lambda_n$ vanishes at $S_{\alpha} = 0$, $\forall \alpha$. However, the global pressure p is not a physical variable, consequently the initial and Dirichlet boundary conditions cannot be directly imposed in the presence of capillary forces.

About numerical methods

Hence, we are left with the (near-)elliptic pressure equation, the parabolic/hyperbolic saturation equation(s). It allows to employ different numerical methods dedicated for these types of equations and solve them separately to reduce the degree of freedom, especially when the flow is multicomponent and miscible. Finally, it is worth pointing out that the pressure equation (2.2.5) must be rewritten in a conservative way in order to apply the Finite Volume Method (FVM).

Being proposed in Chavent (1976), the idea of decoupling the pressure equation is extended to the multicomponent case and used in multiscale methods since it allows to evaluate the velocity field. For decoupled formulations 2.2.1, the sequential solution is

usually applied – IMPES (IMplicit Pressure EXplicit Saturation) method, where the pressure equation is first solved implicitly, the explicit discretization of the saturation equation is then employed with fixed fluxes (similar to sequential modeling single-phase flow and multicomponent transport). The FIM (Fully Implicit Method) is usually employed for the PS formulation that requires to solve the whole system of nonlinear equations simultaneously. The AIM (Adaptive Implicit Method) allows to combine these methods by applying different levels of implicitness to each grid block dynamically in time (Thomas et al., 1983; Russell et al., 1989). The comparative study of three methods was presented in Marcondes et al. (2009) and highlighted the advantages of the AIM.

2.2.2 Compositional flow

When the phase transition is possible, the mass balance is formulated for each component k , $k = 1, \dots, N_s$. So, the compositional flow system consists of N_s partial differential equations

$$\frac{\partial}{\partial t} \sum_{\alpha} \phi S_{\alpha} \rho_{\alpha} X_{\alpha,k} + \nabla \cdot \left(\sum_{\alpha} \rho_{\alpha} X_{\alpha,k} \mathbf{u}_{\alpha} \right) = q_k \quad (2.2.12)$$

for $2N_f + N_f N_s$ natural variables: $2N_f$ phase pressures and saturations, $N_f N_s$ mass fractions. To close the system, we add $N_f - 1$ capillary pressure relations (2.1.5), the total saturation (2.1.1), N_f mass fraction (2.1.3) sums and $(N_f - 1)N_s$ relations of phase equilibria. For instance, assuming $N_f = 2$ and $\alpha, \beta = \{g, l\}$, the natural variables (Coats, 1980) can be taken as the primary variables: p_g , S_l and $N_s - 2$ mass fractions $X_{g,k}$. While the phase α disappears, the set of variables changes to p_g and $N_s - 1$ mass fractions $X_{\beta,k}$, $\beta \neq \alpha$. The set of phases and their composition are usually defined by the phase stability test (Michelsen, 1982a) and flash calculation (Michelsen, 1982b). The formulations and methods for solving compositional flow will be discussed later in Sec. 3.1.1.

2.2.3 Intermediate case

The black-oil model, aimed at modeling the hydrocarbon system, represents an alternative between immiscible and miscible flow. There are three phases (water, oil, gas) and three components (*water, oil, gas*). The oil-gas interactions are allowed, while the water phase is unreactive with the rest of two phases. As an extension of the black-oil model, the

limited compositional model was developed to eliminates the last restriction. To introduce solubility, vaporization, the phase transfer is expressed by a volume ratio (normalized volume change due to reaction), which consequently appears in the accumulation term. The complete description of three-phase flow for hydrocarbon systems can be found in Peaceman (1977).

2.3 Immiscible two-phase flow model

The present development of multiphase flow module is seen as a preliminary work before coupling with the reactive transport framework HYTEC. The following hypotheses are made:

- the immiscible two-phase flow formulation, $\alpha = \{w, n\}$;
- some amount of water is always present in REV (CHESS numerical limitation: the chemical speciation library needs non-zero quantities of water to avoid infinite concentrations);
- isotropy of the intrinsic permeability, $\mathbb{K} = K\mathbf{I}$;
- isothermal system, no heat conservation equation;
- no pore compressibility $\partial\phi/\partial p = 0$, no geochemical deformation;
- incompressible flow.

Since the wetting phase does not disappear, the PS formulation can be applied, where $p = p_w$ and $S = S_n$ are the primary variables. The flow problem consists of two PDEs on $\Omega \times (0, T)$, $\Omega \in \mathbb{R}^N$, $N = 1, 2, 3$

$$\frac{\partial}{\partial t} \phi(1 - S) \rho_w - \nabla \cdot (\rho_w \lambda_w K (\nabla p - \rho_w \mathbf{g})) = q_w, \quad (2.3.1)$$

$$\frac{\partial}{\partial t} \phi S \rho_n - \nabla \cdot (\rho_n \lambda_n K (\nabla(p + p_c) - \rho_n \mathbf{g})) = q_n \quad (2.3.2)$$

with the boundary conditions

$$\begin{aligned} p(\mathbf{x}, t) &= p_d(\mathbf{x}, t) \text{ on } \Gamma_w^D, & \rho_w \mathbf{u}_w \cdot \mathbf{n} &= \psi_w(\mathbf{x}, t) \text{ on } \Gamma_w^N, \\ S(\mathbf{x}, t) &= S_d(\mathbf{x}, t) \text{ on } \Gamma_n^D, & \rho_n \mathbf{u}_n \cdot \mathbf{n} &= \psi_n(\mathbf{x}, t) \text{ on } \Gamma_n^N \end{aligned} \quad (2.3.3)$$

and with the following initial conditions

$$p(\mathbf{x}, 0) = p_0(\mathbf{x}), \quad S(\mathbf{x}, 0) = S_0(\mathbf{x}), \quad \mathbf{x} \in \Omega. \quad (2.3.4)$$

Under the taken assumptions, porosity, viscosity and density are constant. The equations (2.3.1) and (2.3.2) can be divided by ρ_w and ρ_n , respectively. To close the system, the relative permeability and capillary pressure models should be imposed.

Further developments on the coupling (Ch. 3) will extend this formulation to compressible, viscous and miscible flow. Modeling non-isothermal flow and anisotropy is outside the scope of this work.

2.4 Numerical method

We present the description of numerical methodology chosen for solving the two-phase flow problem, for the PS formulation in particular: first, the discretization in space and time; second, the linearization and Newton's algorithm; next, solving linear system and finally, the Jacobian structure.

2.4.1 Discretization

Volume averaging

The FVM based on Voronoï meshes is employed for Eqs. (2.3.1) and (2.3.2). Let us introduce the following definitions:

Ω_i is the element with the center \mathbf{x}_i such that $\bar{\Omega} = \bigcup_{i \in \{1, N\}} \bar{\Omega}_i$,

$$|\Omega_i| = \int_{\Omega_i} d\omega. \quad (2.4.1)$$

$\Gamma_{ij} = \partial\bar{\Omega}_i \cap \partial\bar{\Omega}_j$ is the interface between the elements Ω_i and Ω_j ,

$$|\Gamma_{ij}| = \int_{\Gamma_{ij}} d\gamma. \quad (2.4.2)$$

$I_i^{int} = \{j : \Gamma_{ij} \neq \emptyset\}$ is the index set of element Ω_j that has the common edge with the element Ω_i .

$I_{\alpha, il}^N = \{l : \Gamma_{il} = \partial\bar{\Omega}_i \cap \Gamma_{\alpha, l}^N \neq \emptyset\}$ is the index set of edges of element Ω_i that lie on the Neumann boundary Γ_{α}^N .

I_{α}^D is the index set of element lying on the Dirichlet boundary Γ_{α}^D .

By applying the divergence (Ostrogradsky's) theorem to the flux term

$$\int_{\Omega_i} \nabla \cdot \mathbf{F}_{\alpha} d\omega = \oint_{\partial\Omega_i} \mathbf{F}_{\alpha} \cdot \mathbf{n} d\gamma, \quad (2.4.3)$$

where the flux $\mathbf{F}_{\alpha} = \rho_{\alpha} \mathbf{u}_{\alpha}$, \mathbf{n} is the outward normal, we discretize the integrals of Eqs. (2.3.1) and (2.3.2) over the element $\Omega_i, \forall i \in I \setminus I_{\alpha}^D$:

$$|\Omega_i| \frac{\partial}{\partial t} \phi_i (1 - S_i) \rho_{w, i} + \sum_{j \in I_{ij}^{int}} |\Gamma_{ij}| F_{w, ij} + \sum_{j \in I_{w, il}^N} |\Gamma_{il}| (\psi_w)_{ij} = |\Omega_i| (q_w)_i, \quad (2.4.4)$$

$$|\Omega_i| \frac{\partial}{\partial t} \phi_i S_i \rho_{n, i} + \sum_{j \in I_{ij}^{int}} |\Gamma_{ij}| F_{n, ij} + \sum_{j \in I_{n, il}^N} |\Gamma_{il}| (\psi_n)_{ij} = |\Omega_i| (q_n)_i, \quad (2.4.5)$$

where F_α is the approximated normal flux $\mathbf{F}_\alpha \cdot \mathbf{n}$. The Neumann conditions are taken into account by integrating the imposed fluxes, while the Dirichlet approximation is given by

$$p_i = p_{d,i} \quad \forall i \in I_w^D, \quad (2.4.6)$$

$$S_i = S_{d,i} \quad \forall i \in I_n^D. \quad (2.4.7)$$

Next, the two point flux approximation (TPFA) is employed for the interior fluxes:

$$F_w = - \left(\rho_w \frac{k_{rw}}{\mu_w} \right)_{ij} K_{ij} \left(\frac{p_j - p_i}{\|\mathbf{x}_j - \mathbf{x}_i\|} - \rho_{w,i} g \right), \quad (2.4.8)$$

$$F_n = - \left(\rho_n \frac{k_{rn}}{\mu_n} \right)_{ij} K_{ij} \left(\frac{p_j - p_i}{\|\mathbf{x}_j - \mathbf{x}_i\|} + \frac{p_{c,j} - p_{c,i}}{\|\mathbf{x}_j - \mathbf{x}_i\|} - \rho_{n,i} g \right), \quad (2.4.9)$$

where $(\cdot)_{ij}$ is the weighting scheme for interface parameters between Ω_i and Ω_j , $g = \mathbf{g} \cdot \mathbf{n}$. The intrinsic permeability at the interface K_{ij} can be evaluated as harmonic or upstream weighting, while for the relative permeability, the upstream treatment is prerequisite

$$k_{r\alpha,ij} = \begin{cases} k_{r\alpha,i} & \text{if } \left(\frac{p_{\alpha,j} - p_{\alpha,i}}{\|\mathbf{x}_j - \mathbf{x}_i\|} - \rho_{\alpha,i} g \right) \geq 0, \\ k_{r\alpha,j} & \text{else.} \end{cases} \quad (2.4.10)$$

Implicit (Backward Euler) in time

The Fully Implicit (Backward Euler) method is used because of its unconditional stability. Assuming $t^n \in (0, T)$ such that

$$0 = t^0 < \dots < t^n < t^{n+1} < \dots < t^M = T, \quad \Delta t^{n+1} = t^{n+1} - t^n, \quad \sum_{n=0, \dots, M} \Delta t^{n+1} = T,$$

and the mass accumulation $A_{\alpha,i} = \phi_i S_{\alpha,i} \rho_{\alpha,i}$, the discretization yields

$$|\Omega_i| \frac{A_{\alpha,i}^{n+1} - A_{\alpha,i}^n}{\Delta t^{n+1}} + \sum_{j \in I_{ij}^{int}} |\Gamma_{ij}| F_\alpha^{n+1} + \sum_{j \in I_{\alpha,il}^N} |\Gamma_{il}| (\psi_\alpha)_{ij}^{n+1} = |\Omega_i| (q_\alpha)_i^{n+1}. \quad (2.4.11)$$

The relative permeability is discretized by implicit upstream treatment (update at every Newton iteration) guaranteeing stability but increasing truncation errors at the same time, as discussed below.

Weighting of nonlinearities, stability

Settari and Aziz (1975), Aziz and Settari (1979), Peaceman (1977) demonstrated the impact of relative permeability approximation at the interface, in space (from midpoint rule

to upstream) and in time (from explicit to implicit) on the accuracy and stability. Blair and Weinaug (1969) presented the numerical results of the implicit method with the implicit and explicit mobilities and highlighted the efficiency of the fully implicit scheme. Despite its unconditional stability, it is not reasonable to use any large time step. Since such stability is associated with numerical dispersion, the effect of smearing the sharp solution appears (in hyperbolic flows), whose degree can be reduced by grid refining, high-order discretization and total-variation-diminishing (TVD) schemes. Allen (1984) showed the numerical capillary diffusion at the Buckley-Leverett problem with upwind flux discretization based on FEM (Finite Element Method). Later, Bastian (1999) compared the implicit and the Crank-Nicolson scheme with upstream weighting of mobilities and also the implicit method with upstream and central differences treatment. We conclude that the fully implicit method with the implicit upstream mobilities can be carefully employed, as in the aimed applications, the capillary forces are indeed present (parabolic case). Moreover, our ultimate goal is to extend the two-phase flow module to multicomponent flow, Ch. 3.

2.4.2 Solving linearized system

Solving the nonlinear system: Newton's method

The discretization Eq. (2.4.11) yields the system of nonlinear algebraic equations

$$\mathbf{F}(\boldsymbol{\zeta}) = \mathbf{0}, \quad (2.4.12)$$

where \mathbf{F} is the residual function with $\dim(\mathbf{F}) = 2N$, $\boldsymbol{\zeta}$ is the vector comprising $2N$ primary variables:

$$\boldsymbol{\zeta} = (p_1, S_1, \dots, p_N, S_N)^T. \quad (2.4.13)$$

The alternative ordering $\boldsymbol{\zeta} = (\mathbf{p}, \mathbf{S})^T$ is also possible. The approximative solution of the differential function \mathbf{F} can be found by Newton's method. Arising from the 1st Taylor polynomial, the estimation of \mathbf{F} at $(k+1)^{\text{th}}$ approximation of root $\boldsymbol{\zeta}^{k+1}$ evolves into

$$\mathbf{F}(\boldsymbol{\zeta}^{k+1}) = \mathbf{F}(\boldsymbol{\zeta}^k) + \mathbf{J}(\boldsymbol{\zeta}^k)(\boldsymbol{\zeta}^{k+1} - \boldsymbol{\zeta}^k), \quad (2.4.14)$$

where \mathbf{J} is the Jacobian of the vector function \mathbf{F} :

$$J_{ij} = \frac{\partial F_i}{\partial \zeta_j}. \quad (2.4.15)$$

Since ζ^{k+1} is the approximative root, the reformulation of Eq. (2.4.14) leads to solving the linearized system

$$\mathbf{J}(\zeta^k)\delta\zeta^{k+1} = -\mathbf{F}(\zeta^k), \quad (2.4.16)$$

where $\delta\zeta^{k+1} = \zeta^{k+1} - \zeta^k$. To summarize, let us present the iterative procedure in Algorithm 1. Once the error tolerance ε_{Nf} , usually $\varepsilon_{Nf} \in [10^{-8}, 10^{-6}]$, and the maximum number of iterations k_{max} are imposed, we deal with the linearized problem to find $\delta\zeta^{k+1}$, line 5. The iterative method continues until the norm of the residual function is small enough, line 4. If $k > k_{max}^{fl}$, the time step Δt^{n+1} in Eq. (2.4.11) is reduced and Algorithm 1 is repeated.

Algorithm 1 Newton's method for Immiscible Two-Phase Flow

```

1:  $\varepsilon_{Nf} = 1 \times 10^{-6}$ 
2:  $k_{max}^{fl} = 9$ 
3:  $k = 0$ 
4: while  $(\|\mathbf{F}(\zeta^k)\|_\infty \geq \varepsilon_{Nf}\|\mathbf{F}(\zeta^0)\|_\infty) \wedge (k \leq k_{max}^{fl})$  do
5:   find  $\delta\zeta^{k+1}$ :  $\mathbf{J}(\zeta^k)\delta\zeta^{k+1} = -\mathbf{F}(\zeta^k)$ 
6:    $\zeta^{k+1} \leftarrow \zeta^k + \delta\zeta^{k+1}$ 
7:    $k \leftarrow k + 1$ 
8: end while
```

Solving the linear system: GMRES method

The linear system (2.4.16) is solved at each Newton iteration. The matrix \mathbf{J} has a non-symmetric sparse structure and becomes larger with increasing N as expected. Hence, the choice of linear solver is of great importance in multiphase modeling inasmuch as it affects the feasibility and the performance of the whole module. The overview of iterative methods for solving the linear systems can be found in Barrett et al. (1994) and Benzi (2002). It is well known that applying a preconditioner improves the condition number of the matrix and therefore, the convergence rate of compatible linear solver. The preconditioning is regularly employed with the Krylov methods; e.g., GMRES (Saad and Schultz, 1986), BiCGSTAB (Van der Vorst, 1992), that are principal solvers for non-symmetric system. Saad (2003) gave an excellent description of the Krylov subspace methods and preconditioning.

The GMRES method with ILU(0) provided by Sparselib++ (Pozo and Remington, 1996) is employed for the system (2.4.16). The tolerance of GMRES ε_{lin} impacts to the convergence rate of Newton's algorithm and it was suggested that $\varepsilon_{lin} \in [10^{-8}, 10^{-5}]$. The restart option can be defined in range $[10^2, 10^4]$ that depends on the condition number of the Jacobian. The split preconditioning is used for ill-conditioned matrix: left diagonal (explicit) and then right ILU, (Saad, 2003, Ch. 9).

Remarks: The two-stage constrained pressure residual (CPR) preconditioner, initially proposed by Wallis et al. (1985), significantly accelerates the calculations of linear systems arising from the fully implicit scheme (Cao et al., 2005; Voskov and Tchelepi, 2012). The algorithm is designed to take into account the types of differential equations and to employ the relevant methods. For instance, the two-phase flow system comprises a near elliptic pressure equation and a near hyperbolic saturation equation (composition equations). After the linearisation, the system is decomposed into subsystems. First, the pressure subsystem is solved by a linear solver, an Algebraic Multigrid solver (Stüben, 1983); the residual of the entire system is then corrected to the obtained pressure solution. Second, a suitable preconditioner, such as ILU(0), is applied to the corrected system.

Analytical Jacobian

The two-phase PS formulation entails the numerical or analytical $2N \times 2N$ Jacobian calculation. The matrix structure is 2×2 block whose dimension corresponds to the number of equations and to the number of primary variables. The Dirichlet conditions (2.4.6) yield trivial blocks.

The analytical estimation is applied in HYTEC. Now, we detail the Jacobian structure. Neglecting the time indices, we derive the diagonal block of incompressible PS problem with the primary variables (p_w, S_n) for Ω_i (2.4.17) and therefore the contribution of i^{th} block to the non-diagonal block j (2.4.18). Since the Jacobian should be invertible matrix, $\mathbf{J}_{2i,2i} \neq 0$ in the diagonal block (2.4.17). Thus, there is a constraint at the extreme value of saturation $S_n \neq 1$, when (p_w, S_n) are primary variables. The restriction $S_w \neq 1$ appears in case of (p_n, S_w) choice by analogy.

The models of relative permeability and capillary pressure are user-defined. The models of fluid properties will be discussed in Ch. 4. Once all property models are selected,

the Jacobian is automatically assembled. The matrix is constructed by gathering the corresponding functions and their analytical derivatives and it is written in the CRS (compressed row storage) format (Poza and Remington, 1996). The nonlinear functions must then be of class $C^1([0, 1])$. Some extensions and redefinitions of fluid-matrix functions and consequently their derivatives should be made at least on $[0, 1 - \varepsilon]$, that is detailed for capillary pressure and relative permeability functions in Appendices A and B, respectively.

$$\begin{aligned}
\mathbf{J}_{2i,2i} &= \Delta t \sum_{j \in I_{ij}^{int}} |\Gamma_{ij}| \left(\rho_w \frac{k_{rw}}{\mu_w} \right)_{ij} K_{ij} \frac{1}{\|\mathbf{x}_j - \mathbf{x}_i\|} \\
\mathbf{J}_{2i,2i+1} &= -|\Omega_i| \phi_i \rho_{w,i} \\
&\quad - \Delta t \sum_{j \in I_{ij}^{int}} |\Gamma_{ij}| \left(\frac{\rho_w}{\mu_w} \right)_{ij} \frac{\partial k_{rw,ij}}{\partial S_i} K_{ij} \left(\frac{p_j - p_i}{\|\mathbf{x}_j - \mathbf{x}_i\|} - \rho_{w,i} g \right) \\
\mathbf{J}_{2i+1,2i} &= \Delta t \sum_{j \in I_{ij}^{int}} |\Gamma_{ij}| \left(\rho_n \frac{k_{rn}}{\mu_n} \right)_{ij} K_{ij} \frac{1}{\|\mathbf{x}_j - \mathbf{x}_i\|} \\
\mathbf{J}_{2i+1,2i+1} &= |\Omega_i| \phi_i \rho_{n,i} \\
&\quad - \Delta t \sum_{j \in I_{ij}^{int}} |\Gamma_{ij}| \left(\frac{\rho_n}{\mu_n} \right)_{ij} \frac{\partial k_{rn,ij}}{\partial S_i} K_{ij} \left(\frac{p_j - p_i}{\|\mathbf{x}_j - \mathbf{x}_i\|} + \frac{p_{c,j} - p_{c,i}}{\|\mathbf{x}_j - \mathbf{x}_i\|} - \rho_{n,i} g \right) \\
&\quad + \Delta t \sum_{j \in I_{ij}^{int}} |\Gamma_{ij}| \left(\rho_n \frac{k_{rn}}{\mu_n} \right)_{ij} K_{ij} \frac{\partial p_{c,i}}{\partial S_i} \frac{1}{\|\mathbf{x}_j - \mathbf{x}_i\|}
\end{aligned} \tag{2.4.17}$$

$$\begin{aligned}
\mathbf{J}_{2i,2j} &= -\Delta t |\Gamma_{ij}| \left(\rho_w \frac{k_{rw}}{\mu_w} \right)_{ij} K_{ij} \frac{1}{\|\mathbf{x}_j - \mathbf{x}_i\|} \\
\mathbf{J}_{2i,2j+1} &= -\Delta t |\Gamma_{ij}| \left(\frac{\rho_w}{\mu_w} \right)_{ij} \frac{\partial k_{rw,ij}}{\partial S_j} K_{ij} \left(\frac{p_j - p_i}{\|\mathbf{x}_j - \mathbf{x}_i\|} - \rho_{w,i} g \right) \\
\mathbf{J}_{2i+1,2j} &= -\Delta t |\Gamma_{ij}| \left(\rho_n \frac{k_{rn}}{\mu_n} \right)_{ij} K_{ij} \frac{1}{\|\mathbf{x}_j - \mathbf{x}_i\|} \\
\mathbf{J}_{2i+1,2j+1} &= -\Delta t |\Gamma_{ij}| \left(\frac{\rho_n}{\mu_n} \right)_{ij} \frac{\partial k_{rn,ij}}{\partial S_j} K_{ij} \left(\frac{p_j - p_i}{\|\mathbf{x}_j - \mathbf{x}_i\|} + \frac{p_{c,j} - p_{c,i}}{\|\mathbf{x}_j - \mathbf{x}_i\|} - \rho_{g,i} g \right) \\
&\quad - \Delta t |\Gamma_{ij}| \left(\rho_n \frac{k_{rn}}{\mu_n} \right)_{ij} K_{ij} \frac{\partial p_{c,j}}{\partial S_j} \frac{1}{\|\mathbf{x}_j - \mathbf{x}_i\|}
\end{aligned} \tag{2.4.18}$$

2.5 Verification

We present the numerical results of a 1D radial parabolic problem with self-similar solution and a 2D hyperbolic problem.

2.5.1 Self-similar solution to two-phase displacement problem

Modeling the problem obeying the self-similarity allows not only to analyze the physical phenomena, intermediate asymptotic behavior (Barenblatt, 1996) but also to verify the numerical code. The invariant solution $\xi = x/a\sqrt{t}$ was initially discovered for the heat equation and then for several (parabolic PDE) infiltration problems and immiscible displacement (Polubarinova-Kochina, 1952; Barenblatt et al., 1972, Ch. 4 and 6). The latter carried out the self-similarity study considering the capillary pressure effects, radial flow, double porosity; afterward, Chen (1988) extended this class of immiscible, incompressible flow problems by including both capillary and gravity terms. The author, in particular, investigated the axisymmetric case taking the capillary forces into account under hypotheses of the saturated, homogeneous, horizontal, infinite reservoir and of the constant rate injection through a well with the infinitesimal radius. By substituting the self-similar variable in Sys. (2.2.1), it is reduced to a 2nd order ODE. Similar results were numerically demonstrated for the mass and energy equations (O’Sullivan, 1981).

Model description

During the workshop *Intercomparison of numerical simulation codes for geologic disposal of CO₂* initiated by Lawrence Berkeley National Laboratory (Pruess et al., 2002), the axisymmetric problem of constant injection in a long, horizontal, reservoir was stated. The self-similar variable is R/\sqrt{t} , where R is the radial distance from injection well and t is time. Adapting the parameters from the workshop, we model the CO₂ injection by means of the built *non-reactive* flow module. (Recall that at this stage, the flow is isothermal, immiscible, incompressible; chemical reactions are excluded.) Table 2.1 presents the 1D radial flow problem statement. CO₂ is injected at the constant rate 100 kg/s into a 100 km long aquifer. The well radius is 0.3 m. The length of grid elements vary from 0.2 m close to the well, to 1 km. The functions of capillary pressure and relative permeabilities are presented in Fig. 2.2 and Fig. 2.3, respectively.

Table 2.1 – Parameters of the axisymmetric problem admitting self-similar variable R/\sqrt{t} .

Initial Conditions		
$\Omega = [0.3, 10^5 + 0.3] \times [0, 100] \text{ [m}^2\text{]}$		
Ω :	$p_{l0} = 120 \text{ [bar]}$	$S_{g0} = 0$
Boundary Conditions		
$\Gamma^N = [(0.3, 0); (0.3, 100)] \text{ [m]}$		
$\Gamma^D = [(10^5 + 0.3, 0); (10^5 + 0.3, 100)] \text{ [m]}$		
Γ^N :	$\psi_l = 0$	$\psi_g = 1/(2\pi 0.3) \text{ [kg/m}^2\text{/s]}$
Γ^D :	$p_l = 120 \text{ [bar]}$	$S_g = 0$
$\partial\Omega \setminus (\Gamma^D \cup \Gamma^N)$:	$\psi_l = 0$	$\psi_g = 0$
Matrix Properties		
Porosity ϕ	0.12	
Permeability K	$10^{-13} \text{ [m}^2\text{]}$	
Fluid-Matrix Properties		
van Genuchten capillary pressure		
Eq. (2.1.8):	$m = 0.457$	$1/\alpha = 19.61 \text{ [kPa]}$
van Genuchten liquid relative permeability		
Eq. (2.1.11):	$m = 0.457$	$S_{lr} = 0.3, S_{gr} = 0$
Brooks-Corey gas relative permeability		
Eq. (2.1.10):	$\lambda = 2$	$S_{lr} = 0.3, S_{gr} = 0.05$
Fluid Properties		
Density	$\rho_l = 1000 \text{ [kg/m}^3\text{]}$	$\rho_g = 666 \text{ [kg/m}^3\text{]}$
Viscosity	$\mu_l = 10^{-3} \text{ [Pa} \cdot \text{s]}$	$\mu_g = 4.8 \times 10^{-5} \text{ [Pa} \cdot \text{s]}$

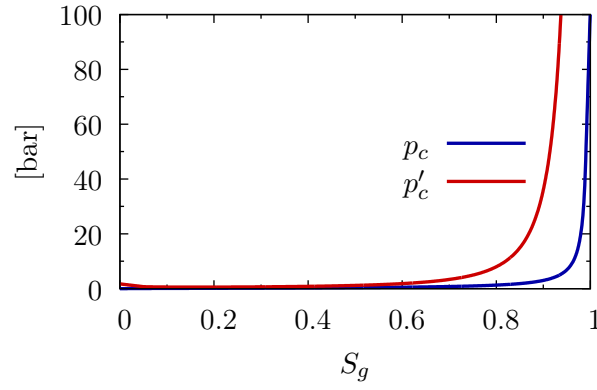


Figure 2.2 – Capillary pressure function of S_g and its derivative: p_c - van Genuchten, Eq. (2.1.8): $m = 0.457$, $1/\alpha = 19.61[kPa]$.

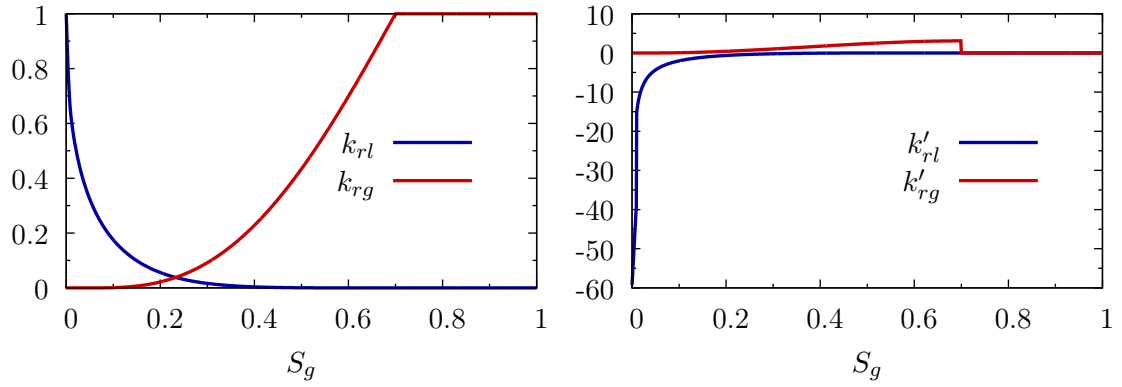


Figure 2.3 – Relative permeability functions of S_g (left) and its derivatives (right). k_{rl} - van Genuchten, Eq. (2.1.11): $m = 0.457$, $S_{lr} = 0.3$. k_{rg} - Brooks-Corey, Eq. (2.1.10): $\lambda = 2$, $S_{lr} = 0.3$, $S_{gr} = 0.05$.

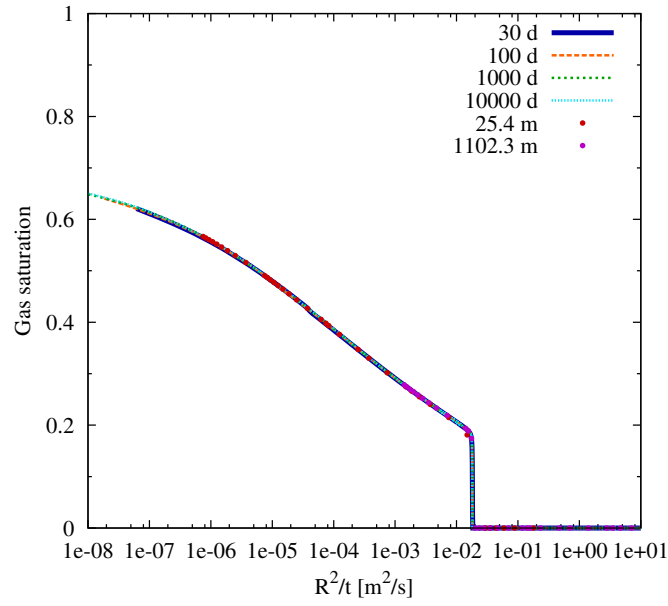
Results

The numerical solution of gas saturation S_g is plotted as a function of the self-similar variable R^2/t in Fig. 2.4(a). The results are presented for the fixed time (30, 10^2 , 10^3 , 10^4 d) and for the fixed radial distance (25.8, 1100.3 m). The numerical solution approximately admits the invariant variable. The time step varies from 5 s to the imposed maximum, 50 d.

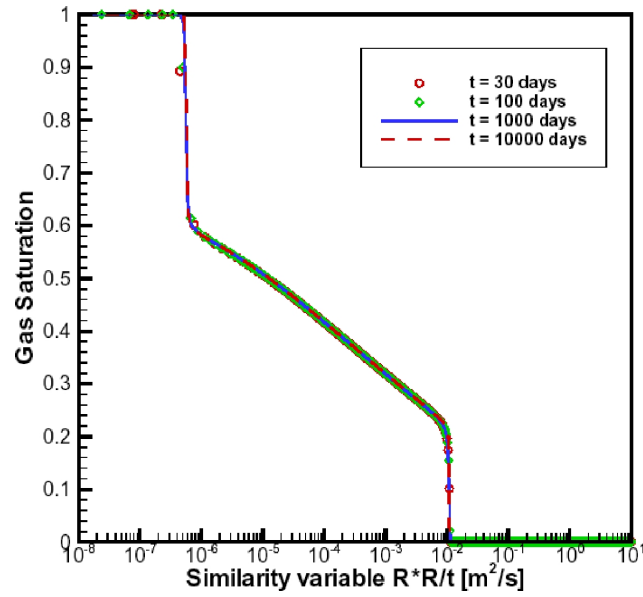
Even taking into account the modifications of the problem (no evaporation of the residual water), the results obtained by HYTEC are in a good agreement with those of TOUGH2-ECO2, Fig. 2.4(b). We detail the differences below. The saturation profile modeled by HYTEC has two regions:

- $R^2/t < 2 \times 10^{-2} [m^2/s]$ that indicates two-phase regime and
- $R^2/t \geq 2 \times 10^{-2} [m^2/s]$ the liquid phase zone,

whereas the third region, $R^2/t \leq 5 \times 10^{-7} [m^2/s]$, Fig. 2.4(b) corresponds to the dry-out zone. Fig. 2.5 displays the time evolution of gas saturation of both HYTEC and of TOUGH2-ECO2 during 10000 d. The apparent shift of saturation profile takes place due to the neglected effects of solubility, fluid and rock compressibility, viscosity, whose role will be shown in Sec. 3.2. Nevertheless, the similarity is preserved and the solution is within the range of the results given by other codes (Fig. 2.6), Pruess et al. (2004). Moreover, the HYTEC's profile is close to that of GEM simulator (ARC in the Fig. 2.6), which also lacks drying-out capabilities.



(a)



(b)

Figure 2.4 – 1D axisymmetric problem: gas saturation S_g as a function of R^2/t . Results of (a) HYTEC and (b) TOUGH2-ECO2 (Pruess et al., 2002). The dry-out zone ($R^2/t \leq 5 \times 10^{-7}$) is not simulated by HYTEC.

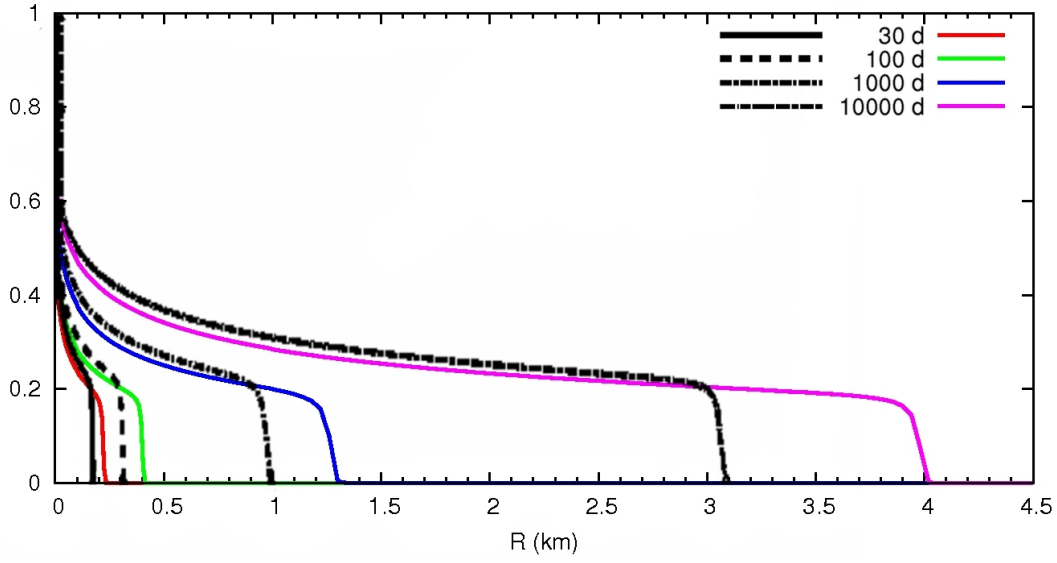


Figure 2.5 – 1D axisymmetric problem: gas saturation S_g . Results of HYTEC (in color) and TOUGH2-ECO2 (Pruess et al., 2002) (black).

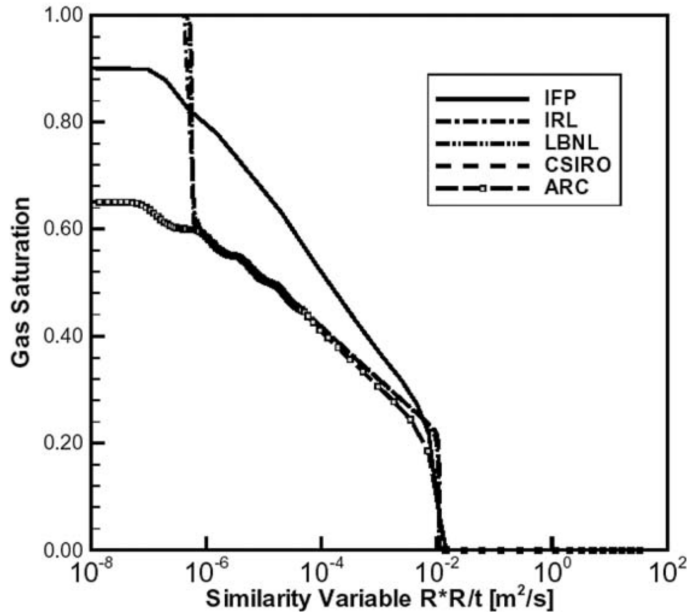


Figure 2.6 – 1D axisymmetric problem: gas saturation S_g as a function of R^2/t . Results of the workshop *Intercomparison of numerical simulation codes for geologic disposal of CO_2* (Pruess et al., 2004): IFP – Institut Français du Pétrole, SIMUSCOPP; IRL – Industrial Research Limited, an in-house version of TOUGH2 and CHEM-TOUGH; LBNL – Lawrence Berkeley National Laboratory, TOUGH2/ECO2; CSIRO – CSIRO Petroleum, an in-house version of TOUGH2/ECO2; ARC – Alberta Research Council, GEM.

2.5.2 Hyperbolic problem. Homogeneous media

By neglecting capillarity, the two-phase system results in the hyperbolic equation which transforms to a Riemann problem under discontinuous initial conditions. The solution is therefore analyzed by deriving characteristic families in order to determine the wave type – rarefaction, shock. The latter forms the discontinuity of solution, that can introduce numerical instability and oscillations. As was mentioned in Sec. 2.4.1, the chosen numerical method smothers the steep solutions. However, it must converge to the physical meaningful solution. The Riemann problem was extensively studied in gas thermodynamics. The detailed physical explanation of phenomena can be found in Loitsyansky (1950, Ch. 4) and Landau and Lifshitz (1986, Ch. 9-11); the mathematical and numerical study was given in Samarskii and Popov (1992, Ch. 1 and 5) and LeVeque (2002). To solve the numerical difficulties of Riemann problem, the Godunov method was proposed that is actually the integrated finite difference or finite volume method. Therefore, applying the FVM allows to model the hyperbolic two-phase flow problems.

In the context of two-phase flow, we refer to the survey by Bastian (1999, Sec. 2.4.2), since the velocity (that becomes a fractional flow function in two-phase flow) depends on the mobility ratio ($M = \lambda_w/\lambda_n$ for waterflooding case), and therefore on the relative permeability models and viscosity ratio. The characteristics analysis yields that the rarefaction and shock wave appearance is subject to the mobility ratio and the inflection point of fractional flow.

Model description

The model of waterflooding process in a 2D homogeneous horizontal reservoir was proposed by Bastian (1999, Sec. 7.2.1). The capillary pressure and gravity are then neglected. Fig. 2.7(a) shows the model geometry. The water (wetting phase) is injected in the left bottom well during 750 d and the host oil (non-wetting phase) is produced from the right top corner. Table 2.2 gives the problem statement; the curves of relative permeability are plotted in Fig. 2.8.

By transforming the notion of M , the viscosity ratio μ is defined as the ratio of the displaced phase viscosity to the injected one: $\mu = \mu_n/\mu_w = 20$ when the water is injected in more viscous fluid. Applying the characteristic method reveals that the solution possesses

different wave states, the rarefaction zone turns to the shock through the tangential point of the fractional flow, see Bastian (1999, Sec. 2.4.2). Hence, reducing the viscosity ratio, for ex. to $1/20$, provides the gradient decrease of rarefaction zone, the solution is steeper, more shock-like than that in case of $\mu = 20$. So, the problem of $\mu = 1/20$, is stated with the same parameters as those of case with $\mu = 20$.

Table 2.2 – Parameters of the hyperbolic problem.

Initial Conditions	
$\Omega = [0, 300]^2 [m]$	
$\mu = 20$	$\mu = 1/20$
Ω : $p_{l0} = 1 [bar], S_{n0} = 0.999$	$p_{l0} = 1 [bar], S_{n0} = 0$
Boundary Conditions	
$\Gamma^N = [(0, 0); (15, 0)] \cup [(0, 0); (0, 15)] [m]$	
$\Gamma^D = [(285, 300); (300, 300)] \cup [(300, 285); (300, 300)] [m]$	
$\mu = 20$	$\mu = 1/20$
Γ^N : $\psi_w = 0.096/\ \Gamma^N\ , \psi_n = 0 [kg/m^2/s]$	$\psi_w = 0, \psi_n = 0.096/\ \Gamma^N\ [kg/m^2/s]$
Γ^D : $p_w = 1 [bar], S_n = 0.999$	$p_w = 1 [bar], S_n = 0$
$\partial\Omega \setminus (\Gamma^D \cup \Gamma^N)$: $\psi_w = 0, \psi_n = 0$	$\psi_w = 0, \psi_n = 0$
Matrix Properties	
Porosity ϕ	0.2
Permeability K	$10^{-10} [m^2]$
Fluid-Matrix Properties	
Capillary pressure: $p_c = 0$	
Brooks-Corey relative permeability	
Eq. (2.1.9), Eq. (2.1.10): $\lambda = 2, S_{wr} = 0, S_{nr} = 0$	
Fluid Properties	
Density $\rho_w = 1000 [kg/m^3]$	$\rho_n = 1000 [kg/m^3]$
Viscosity $\mu_w = 10^{-3} [Pa \cdot s]$	$\mu_n = 20 \times 10^{-3} [Pa \cdot s]$

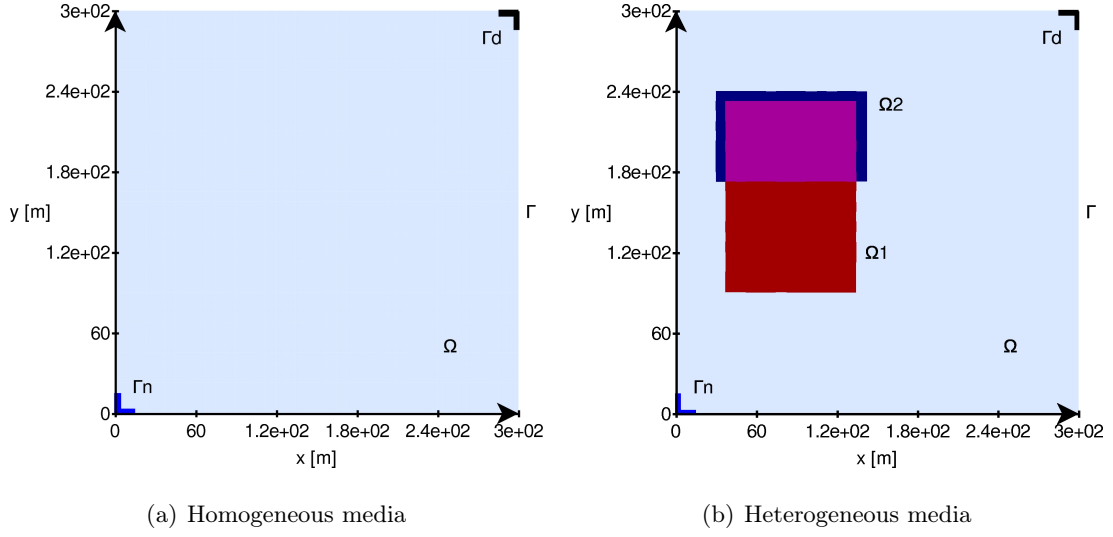


Figure 2.7 – Hyperbolic problem: geometry.

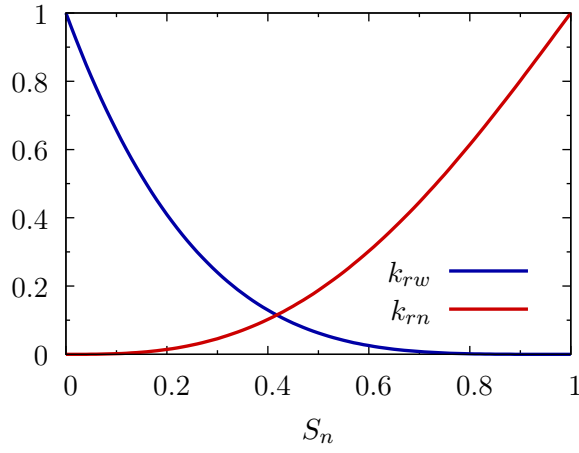


Figure 2.8 – Brooks-Corey relative permeability function of S_n , Eq. (2.1.9), Eq. (2.1.10)
 $\lambda = 2$, $S_{wr} = 0$, $S_{nr} = 0$.

Results

The contour lines of wetting phase saturation S_w for $\mu = 20$ are given on the left column of Fig. 2.9. Refining the grid displays the convergence of the method. As discussed above, the rarefaction zone is followed by the shock front. The PS formulation with primary variable (p_n, S_w) was used in Bastian (1999) and the initial condition is $S_{n0} = 1$, while the chosen formulation PS with pair (p_w, S_n) enforces to adapt it, $S_{n0} = 0.999$. In spite of this difference, the results of UG (Bastian, 1999) and HYTEC are similar, Fig. 2.10.

The case of $\mu = 1/20$ is demonstrated on the right column of Fig. 2.9. Again, the convergence of the method can be illustrated using grid refining. The solution exhibits the gradient decrease – the front sharpening. The injected non-wetting fluid of higher viscosity advances by occupying more volume and the front progress is then slower, comparing with the injection of less viscous wetting fluid.

The method allows to model the rarefaction and shock waves approximately. The convergence and the importance of grid scale were demonstrated. The results are close to those in the reference and provide the physical solution.

2.5.3 Discontinuous permeability

Model description

The model of discontinuous permeability, posed in Bastian (1999, Sec. 7.2.3), points out the numerical difficulties associated with the discontinuity of intrinsic permeability. Based on the hyperbolic problem with homogeneous field Table 2.2, the next discontinuity subzones are added, Fig. 2.7(b):

$$\begin{aligned}\Omega_1 &= [37.5, 135] \times [90, 232.5] [m], \\ \Omega_2 &= [30, 142.5] \times [172.5, 240] [m],\end{aligned}$$

where

$$K = \begin{cases} 10^{-16} & \text{if } \mathbf{x} \in \Omega_1, \\ 10^{-10} & \text{otherwise,} \end{cases} \quad (2.5.1)$$

and

$$S_{w0} = \begin{cases} 0.8 \text{ (0.2)} & \text{if } \mathbf{x} \in \Omega_2 \text{ (and if } \mu = 1/20), \\ S_{w0} & \text{otherwise.} \end{cases} \quad (2.5.2)$$

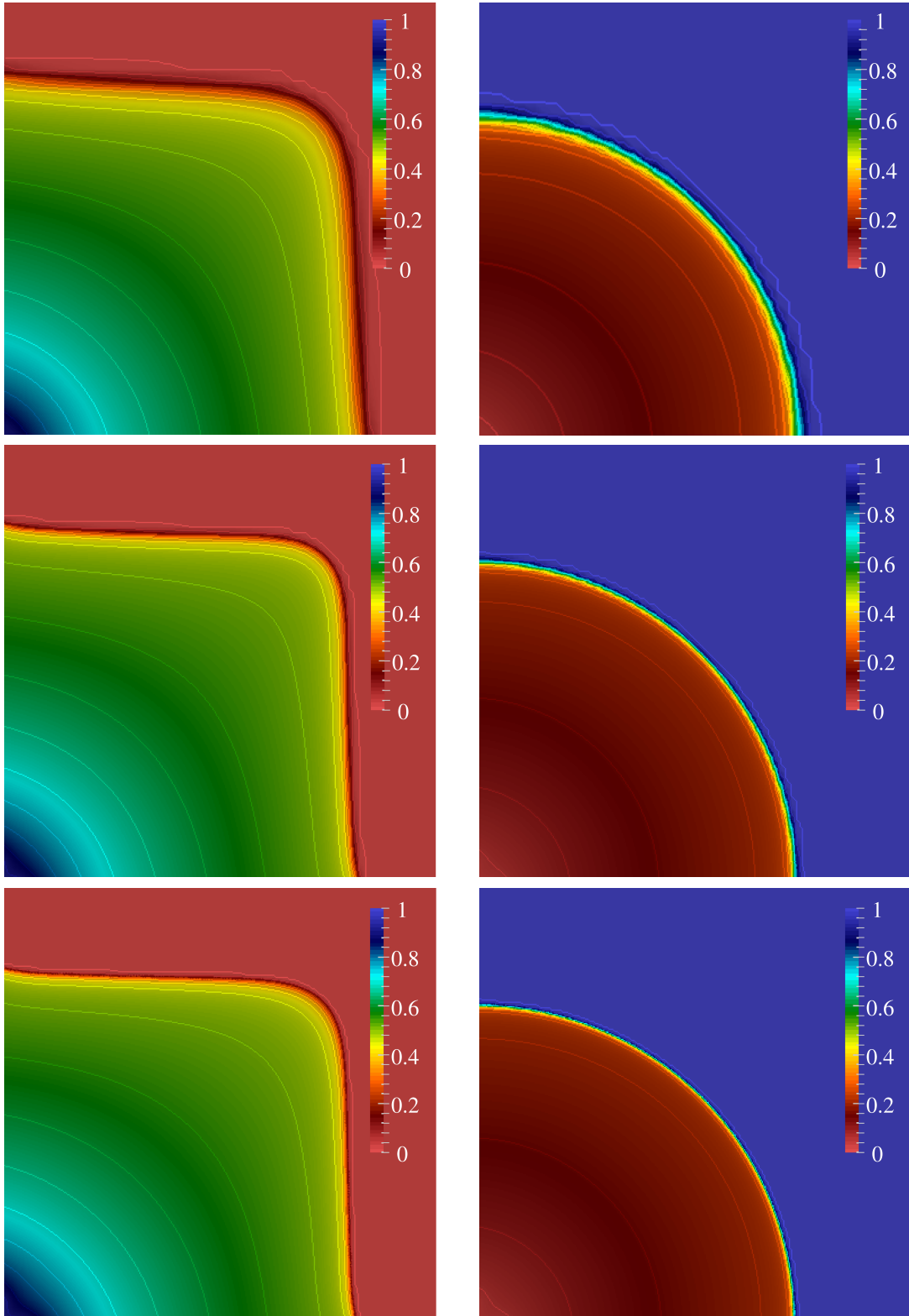


Figure 2.9 – Hyperbolic problem. Homogeneous media. Saturation S_w isolines at 750 d: $\mu = 20$ (left) minimum level - 0.0011, contour interval - 0.05; $\mu = 1/20$ (right) maximum level - 0.9999, contour interval - 0.05. 40×40 (top), 80×80 (center), 160×160 (bottom) elements.

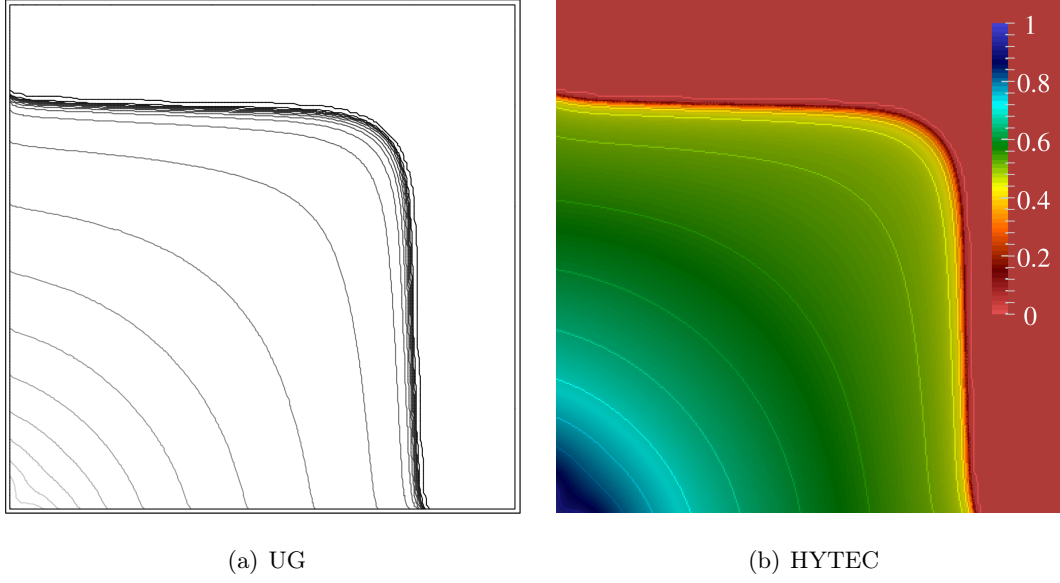


Figure 2.10 – Isolines of saturation S_w , 160×160 elements: (a) results of UG, minimum level - 0.0001, the contour interval - 0.05; (b) results of HYTEC, minimum level - 0.0011, contour interval - 0.05.

Then, the intersection of them $\Omega_3 = \Omega_1 \cap \Omega_2$ presents a low permeability region, containing some amount of injected fluid – 80% saturated. The analogous model is stated for $\mu = 1/20$. Therefore, the initial heterogeneity is provided by the discontinuous saturation and absolute and relative permeability distribution.

Results

Fig. 2.11 shows the results obtained by UG and HYTEC. The contour lines of wetting phase saturation are similar, although there are some differences. HYTEC's front of wetting fluid from $\Omega_2 \setminus \Omega_1$ progresses faster than that of UG, that can be due to the different discretization methods and definition of $\Omega_{1,2}$ ($\Omega_{1,2}$ are slightly shifted comparing with those of Bastian (1999, Sec. 7.2.3)).

For both viscosity ratios, the convergence of method can be seen on Fig. 2.12, where that is expressed by the front sharpening and alignment of Ω_1 , as expected. In water-flooding case $\mu = 20$, the apparent impact of grid refining is additionally highlighted by the stretched form of front. We have seen the nature of solution with different viscosity ratios on the previous model, Sec. 2.5.2. The solution behavior in this model is analogous

to that of homogeneous media.

This test demonstrated a high significance of grid scale, especially in case of high μ , the convergence of the built module and the capability to model the discontinuous permeability.

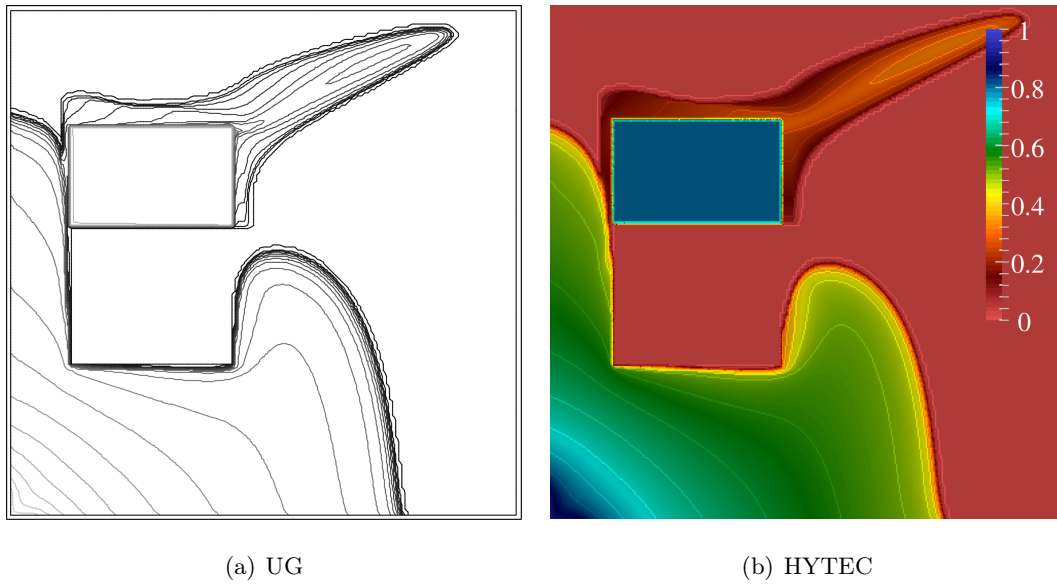


Figure 2.11 – Discontinuous permeability problem. Isolines of wetting phase saturation S_w at 375 d, 160×160 elements: (a) results of UG, minimum level - 0.0001, contour interval - 0.05; (b) results of HYTEC, minimum level - 0.0011, contour interval - 0.05.

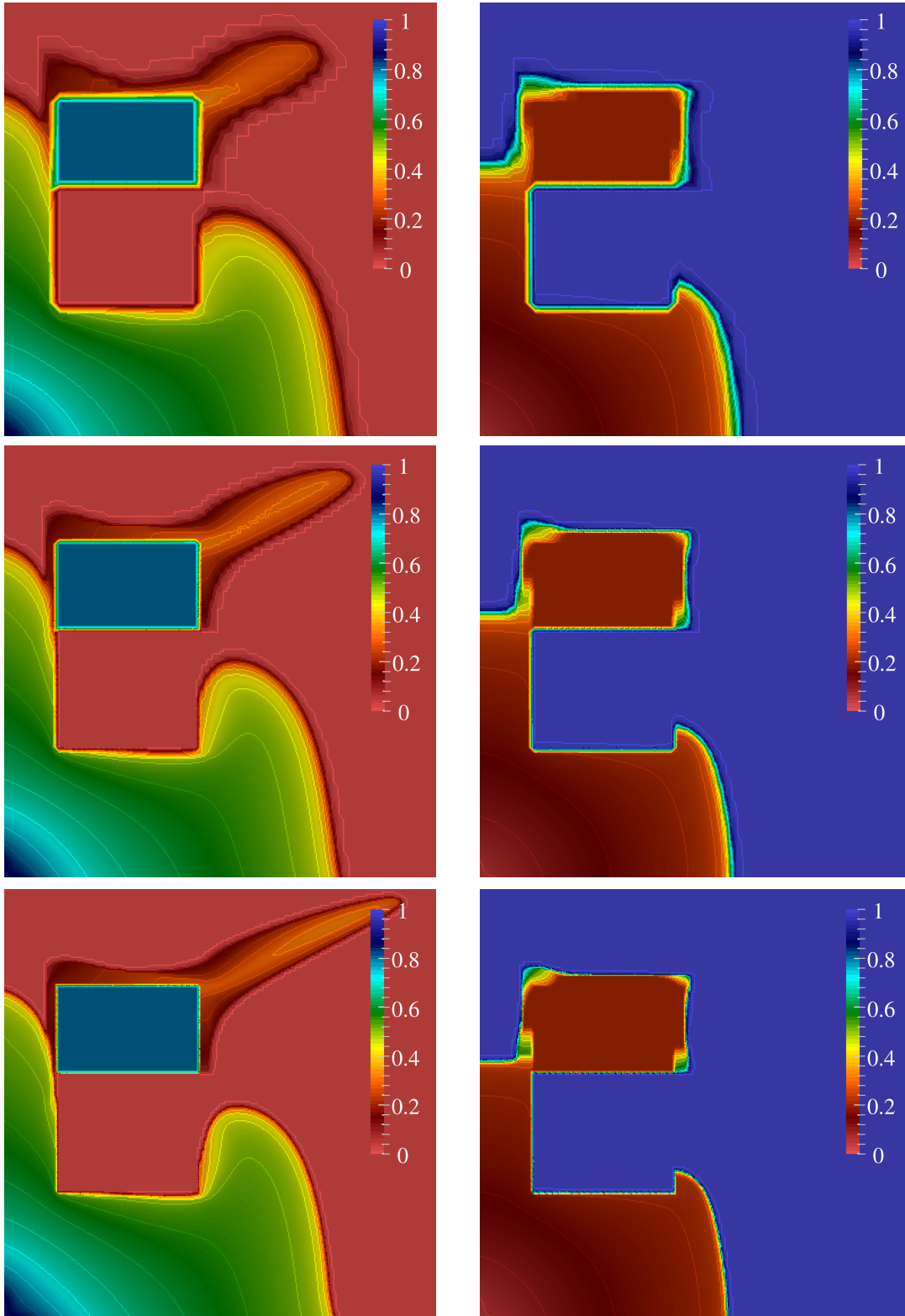


Figure 2.12 – Discontinuous permeability problem. Saturation S_w isolines at 375 d: $\mu = 20$ (left) minimum level - 0.0011, contour interval - 0.05; $\mu = 1/20$ (right) maximum level - 0.9999, contour interval - 0.05. 40×40 (top), 80×80 (center), 160×160 (bottom) elements.

Chapter 3

Coupling Two-Phase Flow and Reactive Transport

Le Chapitre 3 est consacré au couplage de l'écoulement multiphasique multicomposant et du transport réactif. Les formulations, méthodes et problématiques sont discutées pour chaque partie, puis pour leur couplage. Nous proposons une nouvelle approche de la modélisation de l'écoulement multiphasique compressible et son couplage avec le transport réactif, basés sur la séparation d'opérateurs. Puisque l'un des objectifs est de réduire la complexité numérique de la résolution de l'écoulement, la formulation de conservation des phases est employée et couplée au transport compositionnel linéaire et aux équations d'état cubiques. La méthode permet de le découpler comme un module et en conséquence et de l'intégrer dans le cadre de codes de transport réactif. En outre, une approche alternative a été développée pour modéliser séparément le transport multiphasique en utilisant des termes de taux de réaction numériques. Le système couplé est comparé aux codes TOUGH2-ECO2 et DUNE. La vérification de précision repose sur des résultats numériques d'un problème de benchmarking avec une solution auto-similaire. La simulation d'un problème de deux phases et deux composants montre la performance numériques vis-à-vis de la formulation globale implicite. Le code de couplage multiphysique peut intervenir plusieurs processus. Par conséquent, un exercice de benchmarking a été proposé. La chaîne de modélisation de réservoir de gaz naturel a été conçue avec des degrés croissants de complexité pour l'hydrodynamique et la géochimie afin de vérifier et valider les codes de couplage.

Reactive transport simulators aim at modeling complex biogeochemical systems. It is of great importance to be able to extend its abilities to multiphase flow due to its relevance to applications covering a wide range of subsurface activities. This chapter is dedicated to the coupling of the multiphase multicomponent flow (MMF) and reactive transport (RT). The main numerical methods are described for each problem in Sec. 3.1. Sec. 3.2 presents a new coupling approach and the first numerical results. The verification of the chosen methods including a benchmark proposition can be found in Sec. 3.3.

3.1 Overview of multiphase multicomponent flow and reactive transport

The section gives conceptual descriptions of MMF (Sec. 3.1.1) and RT (Sec. 3.1.2) modeling and their mathematical/numerical issues. A list of existing coupling methods is provided in Sec. 3.1.3, before presenting the actual coupling, Sec. 3.2.

3.1.1 Multiphase multicomponent flow

Numerical methods

The formulation of miscible/compositional flow, presented in Sec. 2.2.2, Eq. (2.2.12), involves the mass conservation equation for each component:

$$\frac{\partial}{\partial t} \sum_{\alpha} \phi S_{\alpha} \rho_{\alpha} X_{\alpha,k} + \nabla \cdot \left(\sum_{\alpha} \rho_{\alpha} X_{\alpha,k} \mathbf{u}_{\alpha} \right) = q_k \quad (3.1.1)$$

and results in N_s nonlinear equations. The problem can be solved by FIM or by implicit pressure explicit component (IMPEC) for decoupled formulation, similar to IMPES.

The fully implicit solution involves dividing the compositional problem into the primary and the secondary sets. The equations that need the information of the adjacent cells correspond to the primary set (e.g., the mass balance equations); otherwise, they are associated with the secondary set; e.g., the vapor-liquid equilibria (VLE) equations (that will be discussed below), the saturation, mole/mass fraction constraints. The last set is decoupled and used to eliminate the secondary variables.

By applying the IMPES/IMPEC scheme, the degree of freedom of the primary set is reduced at least twice in comparison with that of FIM, as only the pressure equations

constitute the primary system. At the same time, explicitness imposes time step restriction, arising from the stability analysis. Analogous to the single phase flow, the Courant-Friedrichs-Lewy (CFL) condition for IMPES must be defined:

$$\Delta t \leq \text{CFL} \min \left(\frac{V_{tot}\phi}{u_{in}}, \frac{V_{tot}\phi}{u_{out}} \right) \quad (3.1.2)$$

where $\text{CFL} \leq 1$, \min is the minimum function over all elements Ω_i of the modeled domain, V_{tot} is the volume of the cell, $u_{in/out} = \oint_{\partial\Omega_i} \sum_{\alpha} \mathbf{u}_{\alpha, in/out} \cdot \mathbf{n} \, d\gamma$ is the inflow/outflow rate. Peaceman (1977) and Settari and Aziz (1975) gave the stability analysis and the corresponding CFL condition with regard to the different approximations of relative permeability. Coats et al. (2003) expanded the CFL definition to three phase multicomponent flow. In the advective dominant regime, the restriction can be an important factor if large time steps are required. The AIM combines the advantages of both of them: the method significantly reduces the number of implicit variables, linear solver and total time per Newton iteration and increases the accuracy of solution (Zhou et al., 2011).

Let us conclude, the IMPEC formulation is advantageous in terms of the number of nonlinear equations in the primary set at the cost of the time step restriction, while the FIM proposes the unconditional stability. Here, we have to cite Settari and Aziz (1975):

As the implicitness of the method increases, stability improves, but truncation errors also increase.

Thus, constraints to the saturation, mole/mass fraction and pressure deviations should be imposed. When applying the AIM, the appropriate CFL switching criteria are calculated to define the implicit level of each cell (Zhou et al., 2011). The detailed description of IMPEC, FIM can be found in Chen et al. (2006).

Phase equilibria

We briefly recall the phase equilibria calculation, relying on the thermodynamics basics (Firoozabadi, 1999; Michelsen and Mollerup, 2007; Smith and van Ness, 1987). The compositional flow system (3.1.1) has to be closed by the phase equilibria conditions. The equilibrium criterion, derived from the Gibbs-Duhem equation, dictates a minimum of the Gibbs free energy at constant temperature T , pressure P and composition. Consequently, it requires the equality of the chemical potentials of species i in all phases and then the

equality of the fugacities:

$$f_i^v(T, P, \bar{y}) = f_i^l(T, P, \bar{x}), \quad (3.1.3)$$

where \bar{x} and \bar{y} are the set of mole fractions in the liquid and gas phases. We first present the $\varphi - \varphi$ approach for solving the phase equilibria and give a general description of equation of state (EOS). Then the $\gamma - \varphi$ approach and the numerical methods are discussed.

By substituting the fugacity coefficients φ_i^α in both sides of the phase equilibrium equation (3.1.3) for species i in solution ($\varphi - \varphi$ approach), one can derive:

$$y_i P \varphi_i^v = x_i P \varphi_i^l, \quad (3.1.4)$$

where x_i and y_i are the mole fractions in the liquid and vapor phases, respectively. Eq. (3.1.4) can be rewritten:

$$\ln(\varphi_i^v y_i) = \ln \frac{f_i^v}{P} = \ln \frac{f_i^l}{P} = \ln(\varphi_i^l x_i), \quad (3.1.5)$$

where the fugacity coefficients can be calculated by EOS.

An EOS establishes a relation between pressure, molar volume and temperature (PvT) and allows to evaluate fugacity, solubility, enthalpy and density. For any pure homogeneous fluid in the equilibrium state and for the region of single phase, this relation can be expressed by $f(P, v, T) = 0$. The simplest one is the ideal gas law:

$$Pv = RT. \quad (3.1.6)$$

The compressibility factor, defined as $Z = Pv/RT$, is equal to unity if the gas is ideal. Otherwise, if the gas is real, the models considering the intermolecular forces are used. In 1873, J.D. van der Waals proposed the cubic EOS, including the repulsive and attractive terms: $P = P_R + P_A$, that can represent VLE for pure components and mixtures. In order to improve the volumetric properties near the critical point, the attractive part was modified by Redlich and Kwong in 1949 (RK) and it became temperature dependent through the α function appeared in P_A . Pursuing the idea to better predict the VLE, the α function was elaborated by Soave in 1972 (SRK). Furthermore, Peng and Robinson (PR76) modified the attraction pressure P_A to attain a better estimation of critical compressibility factor Z , liquid density and vapor pressure in comparison with those of SRK.

The cubic EOS can be reformulated as a cubic equation on Z , that has three roots for a given P , two of them can be complex. Depending on the number of phases, the real roots

correspond to the Z in the liquid or vapor phases. Note that the EOS is separately solved for liquid and vapor phases by using the set \bar{x} and \bar{y} , respectively. The cubic equation can be solved for v either analytically or numerically (by an iterative method). Once the roots are found, the calculation of fugacity coefficients $\ln \varphi_i^\alpha$ of a pure component or of a component in a mixture is straightforward by the EOS. The latter is performed by means of the binary interaction parameters and the mixing rules (Peng and Robinson, 1976; Firoozabadi, 1999; Smith and van Ness, 1987; Michelsen and Mollerup, 2007). Another advantage of EOS is the unique way of PvT properties and VLE calculation even in the critical region. The fully implicit solution of the EOS compositional flow was proposed in the pioneer work of Coats (1980).

The EOS equations cannot embrace the wide range of chemical systems, the activity models are then employed to describe the (real) behavior of species in solution. Therefore, the RHS of VLE Eq. (3.1.4) is expressed by applying either the symmetric or the asymmetric activity coefficient γ ($\gamma - \varphi$ approach). The latter uses the reference state in a solvent. Denoting the symmetric activity coefficient of species i at infinite dilution in a solvent k by γ_i^∞ such that $\gamma_i^\infty = \lim_{x_i \rightarrow 0} \gamma_i$, $i \neq k$ yields

$$f_i^l = K_i^{h,\infty} \tilde{\gamma}_i x_i, \quad (3.1.7)$$

where $\tilde{\gamma}_i = \gamma_i / \gamma_i^\infty$ is the asymmetric activity coefficient. Hence, it follows by the definition of $\tilde{\gamma}_i$ that

$$\lim_{x_k \rightarrow 1} \tilde{\gamma}_k = 1, \quad (3.1.8)$$

$$\lim_{x_i \rightarrow 0, i \neq k} \tilde{\gamma}_i = 1. \quad (3.1.9)$$

$K_i^{h,\infty}$ comprises the Henry's law constant K_i^h , representing the solute-solvent relations, and the Poynting factor, providing the pressure effect on K_i^h :

$$\left(\frac{\partial \ln K_i^h}{\partial P} \right)_T = \frac{\bar{v}_i^\infty(T, P)}{RT}, \quad (3.1.10)$$

where $\bar{v}_i^\infty(T, P)$ is the partial molar volume of species i at the infinite dilution in the solvent k (e.g., water). Both approaches ($\varphi - \varphi$ and $\gamma - \varphi$) are connected: $\gamma_i = \phi_i / \phi_0$ at given T and P , where ϕ_0 is the fugacity coefficient at a reference state. The detailed derivation can be found in Michelsen and Mollerup (2007).

The phase equilibria can be performed by flash calculation. The K-value/equilibrium ratio is defined as $K_i = y_i/x_i$. By applying either $\varphi - \varphi$ or $\varphi - \gamma$ approach, one derives

$$K_i = \frac{\varphi_i^l}{\varphi_i^v} = \frac{K_i^{h,\infty} \tilde{\gamma}_i}{P \varphi_i^v}. \quad (3.1.11)$$

The K-values can be calculated from the EOS, activity models, simplified and tabulated as PT dependent coefficients. Under the assumption of ideal mixture, the K-values are composition independent.

Supposing that V and L are the phase mole fractions, $V + L = 1$ and that

$$z_i = x_i L + y_i V, \quad (3.1.12)$$

where z_i is the overall mole fraction of species i , the fractions can be expressed as

$$x_i = \frac{z_i}{1 + V(K_i - 1)}, \quad (3.1.13)$$

$$y_i = \frac{z_i K_i}{1 + V(K_i - 1)}. \quad (3.1.14)$$

The constraint on mole fractions

$$\sum_{i=1}^{N_s} (y_i - x_i) = 0 \quad (3.1.15)$$

evolves into the Rachford-Rice equation (Smith and van Ness, 1987)

$$F(V) = \sum_i^{N_s} \frac{z_i (K_i - 1)}{1 + V(K_i - 1)} = 0. \quad (3.1.16)$$

There are $2N_s + 1$ unknowns, V (or L), \mathbf{x} and \mathbf{y} and the system of $2N_s + 1$ equations, Eqs. (3.1.3), (3.1.12) and (3.1.15). The successive substitution (SS) method and Newton's method are usually employed to solve the system. At each SS iteration, we solve the Rachford-Rice equation on V at given P , T and \mathbf{z} by Newton's method; then the mole fractions are calculated by Eqs. (3.1.16), (3.1.13) and (3.1.14); the K-values are updated through Eq. (3.1.3). When the estimate is close to the solution, Newton's method is applied to the system, Eqs. (3.1.3), (3.1.12) and (3.1.15), that accelerates the convergence. The analysis of the Rachford-Rice function, its derivative, the convergence and the stability of flash calculation are given in Firoozabadi (1999), Michelsen and Mollerup (2007). Different approaches of the flash calculation exist but the subject is outside the scope of this study.

Solving phase equilibria can be either integrated in the Jacobian of compositional flow system or separately coupled with the conservation system. For each cell of single phase, the phase stability test (Michelsen, 1982a) is performed to capture the switch to different phase region, single or two-phase. Then, for each cell of two-phase, the phase composition is defined through the thermodynamic flash.

The important difference between using the EOS models and flash is the way of calculation: the majority of EOS models can be performed analytically, while the flash requires the iterative procedure if a real mixture is considered. Note that the EOS approach is equivalent to the K-values definition and one iteration of flash calculation.

Primary variables

Solving the compositional flow system (3.1.1) by FIM or IMPEC, several formulations of primary variables can be applied to deal with the phase (dis)appearance. The commonly used technique is to choose the natural variables (NVF), pressures, saturations, mole fractions, then the primary variable switching is applied (Coats, 1980): the variables are changed according to the present phases, that is defined by means of the phase stability test. Different combinations are also employed; e.g., pressure (water), saturation(s) and depending on the system state, liquid or gas mole fractions, $S_g y_i$ or $S_l x_i$ with the constraint equations (on saturation and on mole fractions), V or L, z_i , $\ln K_i$, for an overview of molar and volume formulations see Voskov and Tchelepi (2012).

To avoid the variable switching, several alternative choices of fixed primary variables were proposed, on the example of two-phase $\{g, l\}$ two-component $k = \{w, n\}$ or/and compositional problem.

- Gas pressure and total concentrations through the concept of extended saturation: Abadpour and Panfilov (2009) extended the gas saturation out of range $[0, 1]$ that indicates single-phase fluid, the authors demonstrated the equivalence of the two-phase and pseudo two-phase formulations.

Remarks: The widely used negative flash method (Whitson and Michelsen, 1989) allows to calculate the gas phase molar fraction. Its value can also be out of range $[0, 1]$ and indicates the number of phases.

- Liquid pressure and component mass: Acs et al. (1985) proposed a general purpose

compositional formulation by choosing one pressure and the component masses. The system of N_s components comprises a volume balance equation and the N_s mass balance equations.

- Liquid pressure and mass concentration/density of component n in the liquid phase $\rho_{l,n} = \rho_l X_{l,n}$ (two-phase two-component problem): the reformulation of solubility or molar fraction of non-wetting component in the liquid phase as a function of pressure using Henry's law was proposed (Ippisch, 2003). Bourgeat et al. (2009) used the Henry's law to impose a criteria of phase (dis)appearance, $\rho_{l,n} = \rho_{l,n}(p_g) = \rho_{l,n}(p_l + p_c(S_g))$, so then $S_g = S_g(\rho_{l,n}, p_l)$ using the inverse capillary function.
- Liquid pressure, liquid saturation and $X_{l,n}$ (two-phase two-component problem): Jaffré and Sboui (2010) applied the phase transition criteria of Bourgeat et al. (2009) in the problem system as nonlinear complementarity constraints that implies the third variable.
- Gas pressure and liquid pressure (two-phase two-component problem): Angelini et al. (2011) chose p_g instead of $\rho_{l,n}$, also making use of the capillary pressure extension and the solubility function.
- Capillary pressure and gas pressure (two-phase two-component problem): Neumann et al. (2013) also employed the inverse capillary pressure and the relation $x_{l,n} = x_{l,n}(p_g)$ but with the nonlinear solubility function.
- Pressures, saturations, fugacities (PSF): Lauser et al. (2011) proposed the extended but fixed set of primary variables, pressures, saturations, fugacities, with the complementarity condition, solving the problem by the semi-smooth Newton method; the method avoids the flash calculation.
- Tie-line based (gamma) variables: Entov et al. (2002) demonstrated that the thermodynamic equilibrium problem could be parametrized by projecting the compositional system on the tie-line space Γ . Using this compositional space parametrization (CSP) approach, Voskov and Tchelepi (2009) introduced a preprocessing step that provides tabulated solutions on the tie-lines of interest. The composition is linearly interpolated within the simplex if it lies close to one of the known tie-lines; otherwise,

an equation of state based procedure is applied. The compositional space adaptive tabulation method can be employed in order to avoid the phase stability test and accelerate/replace the flash calculations: the tie-lines are adaptively constructed, the Γ -space is discretized by Delaunay tessellation and the composition is interpolated inside the simplex.

Zaydullin et al. (2012) have presented an adaptive CSP method, using a new set of N_s variables for subcritical region: pressure, liquid phase volume fraction and tie-line parameters γ . These fixed variables, called gamma variables, are switched to pressure and overall composition in the supercritical space. The new gamma variables entail a different interpolation of the composition within the simplex and a modified flash calculations without the Rachford-Rice equation (Smith and van Ness, 1987).

Masson et al. (2014b) compared the NVF (Coats, 1980), PSF (Lauser et al., 2011) and extended pressure-pressure formulation (PPF) (Angelini, 2010) formulations on the numerical convergence in the context of phase (dis)appearance. The authors demonstrated the equivalence of formulations and revealed the similarly better results of NVF and PSF (in terms of the convergence) than those of PPF, especially modeling the gas appearance by transport. Ben Gharbia et al. (2015) presented the NVF and PPF performance on the multicomponent system. The authors showed that the numerical difficulties of PPF appearing in the transition zones caused the time step to decrease and then made it less efficient than the NVF; however, the nature of this issue was not defined in the work.

By applying the flash calculation, described above, one can solve the system (3.1.1) on P and the overall composition \mathbf{z} , Eq. (3.1.12), that are consistent during the phase transition. The comparison between the overall composition and NVF was given by Lu et al. (2010), where the formulation using the total mole fractions displayed a higher convergence over the NVF formulation for CO_2 injection problem. Voskov and Tchelepi (2012) provided a comparative study of the NVF, molar and volume formulations and illustrated that the NVF is a preferable choice for isothermal immiscible and near-miscible problems of gas displacement in heterogeneous medium, especially when the phase state changes.

The implicit methods, especially AIM, are efficient methods for solving the compositional problem of the advective dominant flow. Despite the changing of the Jacobian

structure, the NVF (Coats, 1980) is still one of the best performing and most reliable approaches. However, the PSF and the overall composition formulations have recently demonstrated a good convergence in Masson et al. (2014b) and Lu et al. (2010), respectively, and need more investigation.

3.1.2 Reactive transport

The numerical methods of RT (Yeh and Tripathi, 1989, 1991; Steefel and MacQuarrie, 1996; Saaltink et al., 2001; de Dieuleveult et al., 2009) are divided into the operator splitting (OS) and global implicit approach (GIA): the former reduces the degree of freedom by splitting the RT into the transport and chemistry modules and by solving them sequentially (non-)iteratively, while the latter handles the full system implicitly. Omitting the source terms, porosity and variable saturation, the reactive transport system can be presented as

$$\frac{dT}{dt} = \mathfrak{T}(C), \quad (3.1.17)$$

$$\mathfrak{R}(X, T) = 0, \quad (3.1.18)$$

where the vectors T and C are the total and mobile total concentration vectors and are supposed to be transposed in the transport block (3.1.17), \mathfrak{T} is the transposed transport operator vector, $\mathfrak{R}(X, T)$ is the chemistry block solved on the unknowns X (e.g., basis species) with the input total concentration vector T . Once X is known, the total and mobile total concentration vectors T and C can be defined, $S(X) = C$. Next, we point out the concepts of classic approaches from the sequential non-iterative approach (SNIA) to the global differential algebraic equations (DAE) approach.

SNIA

Assuming that the fluid is first transported and the reactions occur just after the displacement, the transport and chemistry blocks are separately (by OS) and sequentially solved. The scheme of SNIA for the time step Δt^{n+1} is expressed as

$$\begin{aligned} T^{tr} &= T^n + \Delta t^{n+1} \mathfrak{T}(C^n), \\ \mathfrak{R}(X^{n+1}, T^{tr}) &= 0, \end{aligned} \quad (3.1.19)$$

where the explicit time discretization is employed for the transport, then the chemistry module is applied for each cell separately. It is well known that the SNIA is associated

with the persistent error for the decay problem that was demonstrated by Valocchi and Malmstead (1992). To increase the accuracy, one can make use of the strang splitting approach, proposed by Strang (1968). The whole problem is divided into three parts: the advective transport for the half time step, then the chemistry step and finally, the dispersive transport for the second half time step.

SIA

The sequential iterative approach (SIA) allows to preserve the OSA and to solve the system globally by Picard's method. The implicit scheme is applied for the transport and then the chemistry module is used for each cell as in SNIA. Therefore, the nonlinear system arising from the chemical operator is solved inside Picard's method. The SIA scheme can be written as

$$\begin{aligned} T^{n+1,2k+1} &= T^n + \Delta t^{n+1} \mathfrak{T}(C^{n+1,2k+1}), \\ \Re(X^{n+1,2k+2}, T^{n+1,2k+1}) &= 0. \end{aligned} \quad (3.1.20)$$

The transport and chemistry modules iterate until the convergence, that reduces the errors in comparison with those of SNIA (Carrayrou et al., 2004).

DSA

In the global direct substitution approach (DSA), the RT problem is solved for the chemistry variables X , substituted in the transport, that leads to

$$(T(X))^{n+1} = (T(X))^n + \Delta t^{n+1} \mathfrak{T}((C(X))^{n+1}). \quad (3.1.21)$$

We are left with the nonlinear system on X . By applying Newton's method, the derivatives of $T(X)$ and $C(X)$ should be calculated, so the chemistry is incorporated in the transport equations. The DSA system can be reduced by the elimination of mineral reactions at equilibrium (Lichtner, 1985; Saaltink et al., 1998).

DAE

The RT system Eqs. (3.1.17) and (3.1.18) can be generalized as a system of DAE:

$$A \frac{dY}{dt} = B(Y). \quad (3.1.22)$$

A is the block matrix, where the non-zero elements form $A_{11} = I$, $\dim A_{11} = \dim T$, $Y = (T, X, C)^T$ is the unknown vector, $B = (\mathfrak{T}(C), \mathfrak{R}(X, T), S(X) - C)^T$, that is detailed in de Dieuleveult et al. (2009). The RT system (3.1.22) embraces both the transport and the chemistry modules and is then aimed at making use of the accuracy properties of the DAE solvers. Newton's method is globally applied. The size of the final linear system is at least twice as large as that of the OSA and of the DSA, but it is sparser than that of DSA. Carrayrou et al. (2010) displayed the accuracy advantages of DAE solvers.

All presented above methods in Sec. 3.1.2 have pros and cons. The SNIA is computationally efficient due to its structure, although it can provide the consistent mass balance errors, depending on the problem (see above). The SIA retains the modular framework and decreases the splitting errors by iterative procedure; that makes it more computationally expensive than the SNIA by definition and than the DSA with the reduced system for certain problems (Carrayrou et al., 2010). The DSA proposes a more robust method at the cost of solving the nonlinear transport system. Using reduction techniques allows to minimize the system. The DAE method entails the largest system solution for which the DAE solvers are supposed to be employed.

3.1.3 Existing approaches

A list of decoupled and sequential multiphase multicomponent flow (MMF) and reactive transport (RT), provided below, illustrates the diversity of solution schemes.

- CodeBright (Olivella et al., 1996) represents a mass balance for water, air and salt species. This simulator is coupled with solved by DSA/SIA RT, code CHEPROO, (Bea et al., 2009; Saaltink et al., 2004).
- DUNE has recently been extended to RT modeling. The work of Hron et al. (2014) demonstrated the S(N)IA of MMF and RT applied to the growth and transport of microorganisms *Escherichia coli*. Later, in the DuMu^X simulator based on the DUNE framework, Ahusborde et al. (2015); Vostrikov (2014) proposed a sequential non-iterative coupling of two-phase multicomponent flow with RT by separating the whole problem into dominant component flow and minor reactive transport subsystems applying SIA to the last.

- HYDROGEOCHEM, based on FEM, applies a “strong coupling” which consists of iterating saturated/unsaturated (modified Richard’s equation) flow, heat and bio-geochemical transport until convergence is reached (Yeh et al., 2004, 2012). HYDROBIOGEOCHEM provides the reactive transport *under* multiphase flow where the transport equation, called modified hydrologic transport, is adapted to FEM, (Fang, 2003). However, there is no coupling with the multiphase flow.
- iCP (Nardi et al., 2014) is an interface between the multiphysic simulator COMSOL, based on FEM, and the geochemical code PHREEQC (Parkhurst et al., 1999). The (un)saturated flow step is provided by COMSOL, then RT is handled by SNIA where the chemistry is solved in each cell by PHREEQC. The communication between two codes (transfer of updated variables) is user-defined, thus the consistency of coupling is variable. The coupling with two-phase flow is also developed¹.
- IPARS (Wheeler et al., 2012), based on the mixed FEM (MFEM), handles iterative coupling of flow and RT, presented by TRCHEM module. The different flow formulations are available: air-water, oil-water, black-oil, EOS compositional models. The FI, iteratively coupled IMPES/IMPEC can be employed. The multiphase RT are connected with any flow model in IPARS and is time split into advection (Godunov), reaction (second order Runge-Kutta) and diffusion/dispersion (expanded MFEM). The species are divided into those in the flowing phases and in the stationary phases. The fluid and rock properties are composition independent during the RT, nevertheless the fluids and rock can be compressible. During the TRCHEM step, the phase mass transfer does not impact on the flow.

In Peszynska and Sun (2002), the multiphase flow formulation was presented under the hypothesis of slightly compressible fluids and of density and composition independent flow, loosely coupled with the SNI reactive transport divided into advection, reaction and diffusion steps.

- MIN3P (Mayer et al., 2012) combines (un)saturated flow with the RT solved by the direct substitution approach (DSA). MIN3P-Bubble involves the sequential iterative solution of (1) unsaturated flow, (2) reactive transport and (3) gas bubble formation

¹Á. Sáinz, personal communication, 2014.

step that comprises the objective function on the trapped gas saturation, considering that the gas concentrations are expressed by partial pressures through the hypothesis of ideal gas, and finally (4) the relative permeability update (Amos and Mayer, 2006). Furthermore, MIN3P-Dusty embodies the reaction-driven advection, hence, the gas velocity term appears in the transport (Molins and Mayer, 2007).

- MoReS is an EOS and K-value compositional simulator, based on FVM, FIM, IMPES, adaptive implicit method (AIM) are available. The full SNIA of multiphase flow, transport for primary (aqueous) species and chemistry modules is proposed in Farajzadeh et al. (2012) and Wei (2012). In the flow step, the pressure equations and mass conservation for each component are solved, where the components may be hydrocarbons or the total dissolved solids (TDS). The batch reactions are modeled by PHREEQC. The fluid properties depend on the pressure, temperature, the amount of dissolved gases, TDS and salinity. This method is limited to slow chemical reactions and requires small time steps (Wei, 2012).
- NUFT sequentially solves compositional multiphase flow and then the GI system of basis species transport in equilibrium combined with mass balance equations of kinetic reactions (TRANS module) (Hao et al., 2012).
- PFLOTTRAN considers a sequential coupling of compositional flow (water/ CO_2) in MPHASE mode and reactive transport (CHEMISTRY) by applying fully implicit solution of each module (Lichtner et al., 2015; Lu and Lichtner, 2007).
- STOMP chooses a “strong coupling” by iterating flow and SI loop of transport and chemistry. The detailed description is given in White and Oostrom (2006); White et al. (2012). Solving flow system evolves into a nonlinear PDE set for mass conservation of water, air, CO_2 , CH_4 , volatile organic compounds and salt. The geochemical module ECKEChem uses a general paradigm of BIOGEOCHEM for modeling reactive chemicals in batch systems.
- TOUGHREACT (Xu and Pruess, 1998; Xu et al., 2012) introduces the geochemical modeling in TOUGH2 Pruess et al. (1999) by coupling sequentially two-component (water/air) two-phase flow and SI RT. TOUGH2 allows to model different systems (e.g., water/hydrogen) by applying the corresponding EOS module.

- UTCHEM is a chemical-flooding multiphase compositional simulator (Delshad et al., 2000), considering the equilibrium reactions and the polymer/gel and biodegradation kinetics. The compositional flow is carried out by a decoupled formulation which consists of the pressure equation and mass conservation for each component (IMPEC). Then the phase equilibria calculations are performed for the concentrations and saturations and EQBATCH for equilibrium reactions in batch (under hypothesis of ideal aqueous solution).

The following codes are based on GIA to solve fully coupled compositional flow and geochemical reactions.

- AD-GPRS, that initially solved a compositional formulation with molar/natural primary variables by FIM/AIM (Cao, 2002; Zhou et al., 2011), has been extended to geochemical modeling by means of the element-based method (Fan et al., 2012), including the phase equilibria and flash calculations. Both equilibrium and kinetic reactions can be handled.
- COORES implicitly solves the compositional flow by means of the active set method. The active set of variables is established at each iteration, the phase transition is given by flash calculation. The geochemical block is built as a separate module.²
- GEM-GHG provides EOS geochemical compositional modeling (Nghiem et al., 2004). The AIM is employed to reduce the size of nonlinear system. After eliminating the explicit chemical equilibrium rates from mass conservation equations, the set is solved by Newton's method. The neighbour independent processes are also decoupled. Mineral dissolution and precipitation can only be performed as kinetic reactions.
- GPAS initially used FIM for solving EOS compositional flow and was developed for chemical flooding (Pope et al., 2005). The chemistry modeling capabilities were added similar to those of UTCHEM. The chemical species are present only in the aqueous phase. Two approaches are performed: hybrid and fully implicit. The former proposes to solve implicit balance equations for hydrocarbon (like dominant components), then explicit equations for the minor components.

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- STARS applies the different level of implicitness (AIM) to solve the k-value based compositional flow with geochemical reactions, developed by CMG as well as GEM. Mineral reactions are considered kinetic, similar to GEM (Computer Modelling Group, 2009).

3.1.4 Discussion

Both the RT and the MMF formulations had been extensively studied, the development of MMF and RT coupling started later, inheriting the same OSA and GIA as in RT. In the OSA, the general idea is to provide fluxes, pressures and saturations calculated by MMF to RT, similar to the single phase RT modeling. The flow is supposed to be miscible, so the compositional formulation is used, that implies solving the nonlinear system, whose dimension evidently depends on the number of components. Therefore, the efficiency of the method relies on the linear solvers and preconditionners (or/and on the multiscale methods) as well as for the GIA. New methods continue to appear, introducing different levels of coupling between flow and RT, from the weak sequential to the strong SIA. Many variations of coupling approaches exist (Sec. 3.1.3), some works give the method comparisons, suggesting that the coupling issue still needs to be investigated. The detailed conclusion on the methods listed in Sec. 3.1.3, is given in the next section, where we present a new method of integrating the MMF in the RT simulator. In contrast to the existing approaches, the phase formulation is employed to model the multiphase multicomponent system by means of our coupling method (Sec. 3.2), that allows to preserve the flow structure independent of the number of components.

Integrating a compressible multicomponent two-phase flow into existing reactive transport simulator

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Abstract

This work aims at incorporating compressible multiphase flow into the conventional reactive transport framework by an operator splitting approach. A new approach allows to retain the general paradigm of the flow module independent of the geochemical processes and to model complex multiphase chemical systems, conserving the versatile structure of conventional reactive transport. The number of mass conservation nonlinear equations arising from the flow module is minimum, the phase flow formulation is employed. Applying the appropriate equation of state made it possible to precisely describe the compressible multicomponent phases, their thermodynamic properties and relevant fluxes. The proposed flow coupling method was implemented in the reactive transport software HYTEC. The whole framework preserves its flexibility in the further physical and numerical purpose developments. The verification of the coupling is obtained by modeling a problem with a self-similar solution. The simulation of a 2D CO₂ injection problem shows the pertinent physical results and the computational efficiency of the method.

Keywords: Compressible two-phase flow, Reactive transport, Sequential iterative coupling, Operator splitting, HYTEC, EOS

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1. Introduction

1.1. Background/Motivation

Human activity in the subsurface has been expanding and diversifying (waste disposal, mining excavation, high-frequency storage of energy), meanwhile the public and regulatory expectations have been growing. Assessment of each step of underground operations including environmental impact evaluation, relies on elaborate simulators and gives rise to a strong necessity of developing the multiphysics modeling. Reactive transport, a geochemical research and engineering tool, deals with multicomponent systems and sophisticated chemical processes (activity and fugacity correction according to different models, mineral dissolution and precipitation, cation exchange, oxidation and reduction reactions, isotopic fractionation and filiation), in addition to gas evaporation and dissolution [47, 58, 73, 78, 88]. Multiphase flow inherits a strong experience from reservoir engineering researches including thermodynamic modeling of complex phase behavior. In particular, the equations of state were deployed to simulate and study interfacial tension, gas, steam or alkaline injection in oil reservoir and enhanced oil recovery [26, 28, 56, 81].

This work aims at incorporating a compressible multiphase flow module into an existing reactive transport simulator. Our coupling method should therefore meet the following requirements:

1. the new approach should handle the different complex multiphase chemical models and retain the general paradigm of a multiphase flow module independent of the geochemical system conserving the conventional reactive transport structure;
2. the number of mass conservation nonlinear equations arising from the flow module should be minimum such that the reduced flow system preserves the matrix structure in order to minimize the computational intensity;
3. the entire framework should preserve its flexibility towards possible further non-isothermal, geomechanical and domain decomposition developments.

The subject of reactive transport methods was extensively investigated in the last two decades (see below). This work focuses on the coupling

between multicomponent multiphase flow (MMF¹) and reactive transport (RT) modules, starting from the survey of the existing approaches.

1.2. A review of multiphase multicomponent flow and reactive transport codes

1.2.1. Operator Splitting Algorithms between MMF and RT

The strength of the operator-splitting (OS) approach (sequential iterative SIA, or sequential non-iterative SNIA) stems from the framework flexibility allowing to develop and verify each module independently. These are major reasons to choose the OS approach for the coupling between MMF and RT, especially when a hydro-geochemical code is aimed at the extension from the single to two phase flow. The following codes apply the OS approach: CodeBright [57], DuMu^X (based on DUNE) [3, 80], DUNE [33], HYDRO-GEOCHEM (unsaturated) [88, 90], iCP [54], IPARS [60, 83], MIN3P (the bubble model) [47, 51], MoReS [28, 82], NUFT [31], PFLOTTRAN [45, 46], STOMP [85, 84], TOUGHREACT [86, 87], UTCHEM [26], also see [74].

The simulators are chiefly based on the finite volume method (FVM) due to its conservative properties. The general tendency of coupling is first to solve (non-)iteratively flow that provides the velocities. The compositional formulation is usually chosen or alternatively, one solves the conservation equations for the dominant components (*e.g.* water/air). When the two-phase flow system involves two components only (*e.g.* $\{\text{H}_2\text{O}, \text{CO}_2\}$), the final set of equations is obviously identical and reduced to the mass conservation for each component.

Once the flow is established, the RT part can be solved by the OS (SIA, SNIA, predictor-corrector) or global implicit approach (GIA) such as the ordinary differential equations based method (ODE, the chemistry module is used as a black box) or the direct substitution approach (DSA) [70], or the differential algebraic equations based method (DAE) [20], [19]. The comparison of OS and GI approaches applied to RT were given in [12], [20], [70], [75]. In 2001, Saaltink et al. exhibited that SIA was more favorable in case of large grid and low kinetic rates than DSA due to the computer storage increase and slow linear solvers. A decade later, the MoMaS benchmark displayed the reliability of both approaches and the enhanced computational potential of the DSA with the reduction technique [38].

¹The abbreviations are detailed in Table 2.

1.2.2. From Global Implicit Algorithm in RT to Global Implicit Algorithm in MMF&RT

The advantage of GIA is its accuracy at the cost of computational resources. However, this method is becoming more competitive with the increasing computer capabilities and with the advances in the methods that allow to reduce the system of equations. Also, reservoir simulators typically make use of the global formulation for the MMF problem in combination with the fully implicit method (FIM) or the adaptive implicit one (AIM), that unifies the FIM and implicit pressure/explicit concentration (IMPEC). They are therefore naturally extended to the RT using GIA, *e.g.* COORES², GEM-GHG [56], GPAS [62], GPRS [11, 27].

For the global implicit solution of the RT, several techniques propose to reduce the initial mass balance system by linear combination and to eliminate the reaction terms in the equilibrium reactions [38, 69, 50]. Such modifications of the DSA for RT lead to mathematical decoupling of the whole system and consequently make it interesting for global implicit coupling of multiphase flow and reactive transport as was recently demonstrated in works of Saaltink et al. [71] and of Fan et al. [27]. Saaltink et al. proposed a method to introduce the chemistry calculations in conventional multiphase simulator with the idea to keep a minimum number of mass conservation equations taking the fluid phase pressures and porosity as primary variables and expressing all secondary variables as a polynomial function of gas pressure (concentrations of the components, fugacity, pH, salinity etc), that requires a pre-processing to use it for each application. Fan et al. employed the element balance formulation [48] by means of the reduction techniques [38, 50] and the decoupled linearized system on the primary and secondary equations. Saaltink et al. demonstrated that the GIA and OSA gave similar results in the context of CO₂ storage and concluded that the full coupling was not necessary for MMF&RT modeling, from a physical point of view (although the OS can be more computationally expensive).

Meanwhile, Gamazo et al. [29] highlighted the significant impact of geochemical reactions on the phase fluxes by modeling the gypsum dehydration when the non-isothermal flow is chemically restrained, *i.e.* the anhydrite-gypsum paragenesis controls the water activity and hence the evaporation process. The importance of DSA, implicitly connecting the equilibrium het-

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erogeneous reactions and the phase flow, was emphasized in comparison with the decoupled formulation of the flow and the RT: the decoupled formulation overestimates the evaporation, whereas its computational time was reduced by 22.5% (compared to DSA). However, the definition for water activity and liquid density were different for the GI and the OS models. Furthermore, the decoupled flow system comprised the conservation equations for dominant components (water and air), but the coupling between flow and RT was not detailed. Given the nature of the formulations described in this work, it appears likely that using the strong OSA (the SI connection between flow and RT with the precise reaction terms in flow equations) would yield results similar to those of DSA at the cost of additional iterations.

1.3. Preamble to a new approach of MMF&RT coupling

Only dominant components can be considered (the dominant components flow formulation) in order to reduce the number of nonlinear equations arising from the flow system. This is efficient when the impact of the other species are negligible. In geochemical problems, the speciation can vary largely over time and space. Ideally, the dominant components should be adapted locally to preserve the accuracy, but that would require specific treatment for the global flow solution.

If the flow system expands such that many species play significant role in the thermodynamic state and phase displacement, the number of nonlinear equations of compositional flow increases with the number of components, that is computationally expensive.

Despite the reduction techniques, the calculation of the primary nonlinear system is unavoidable. In the geochemical modeling, which encompasses the complex homogeneous and heterogeneous reactions, the primary species usually changes both in space and in time. The system is therefore redefined entailing the modification of Jacobian structure inside Newton's method. Meanwhile, the solution of the transport linear system can be several or even hundred times faster than that of multiphase flow.

Can we then replace the compositional formulation by the phase formulation in order to retain a minimal size of non-linear system? The phase flow coupled to the RT was employed in the work of Peszynska and Sun. Given their problem conditions (slightly compressible fluids and density independent flow), the authors divided the RT into three stages (advection, reaction, diffusion) within the internal time steps by interpolating the fluxes and saturations. Later, Hron et al. simulated *Escherichia coli* growth and transport

under aerobic and anaerobic conditions propounding the sequential coupling of phase flow and the S(N)I RT. The liquid fluid was supposed incompressible, while the gas density depended on phase composition. For advective dominant regime, the transport was solved explicitly in time that led to the time step restriction; CFL = 0.4 and the split transport time step were thus applied. Notice that the same issue arises from the IMPEC scheme where the component equations are solved explicitly.

1.4. Our alternative method

This work proposes a new approach for incorporating the compressible two-phase flow in a conventional reactive transport simulator. It employs the phase flow formulation and preserves all the sustainability and facilities of reactive part. The method inherits the unconditional stability of the fully implicit scheme and can handle the modeling of advective dominant and density driven regimes. The following simplifications are considered in this work: isothermal flow, no geomechanics. The method was applied to the (SIA) reactive transport simulator, HYTEC [41, 78], that was widely evaluated in several benchmarks [22, 12, 40, 76] and in numerous applications, such as cement degradation [24], radioactive waste disposal [25, 23], geological storage of acid gases [18, 35, 39], uranium *in situ* recovery processes [66].

In Sec. 2, a concise description of the governing equations of multicomponent multiphase flow and reactive transport is given. Next, the proposed methods of coupling are detailed in Sec. 3. Sec. 4 then demonstrates the method's applicability and computational performance first on a benchmark problem with a self-similar solution and then, on a 2D CO₂ injection modeling. The conclusions and discussions are finally made in Sec. 5.

Table 1: Nomenclature

Latin Symbols	
a	attraction parameter in Peng-Robinson EOS
a_α	dispersivity in phase α , $[m]$
A_s	specific surface area, $[m^2/m^3 \text{ solution}]$ or $[m^2/kg \text{ mineral}]$
b	van der Waals covolume
c_l^k	total liquid mobile concentration of basis species k , $[mol/kg \text{ } w]$
c_s^k	immobile concentration of basis species k , $[mol/kg \text{ } w]$
c_g^m	gas concentration of basis species m , $[mol/m^3]$

Table 1 Continued: Nomenclature

C_i	concentration of primary species i in chemical module
d	dissolution parameter of transport model
D_α	molecular diffusion coefficient of phase α , $[m^2/s]$
D_α^e	effective diffusion coefficient of phase α , $[m^2/s]$
e	evaporation parameter of transport model
\mathbf{F}	residual function
\mathbf{g}	gravitational acceleration vector, $[m/s^2]$
\mathbf{J}	Jacobian
k	kinetic constant, $[mol/m^2/s]$
k_{max}^{fl}	maximum number of iterations in flow coupling
$k_{r\alpha}$	relative permeability of phase α
\mathbb{K}	intrinsic permeability tensor, $[m^2]$
K_i	K-value/equilibrium ratio
K_j	equilibrium constant of reaction j
K_s	solubility constant of solid phase
K_i^h	Henry's law constant
M	molecular weight, $[kg/mol]$
\max	maximum over the modeled domain
n_α	quantity of matter in phase α
\mathbf{n}	normal vector
N_f	number of fluid phases α
N_g	number of gas species
N_c	number of basis species in chemistry module
N_p	number of phases
N_r	number of independent chemical reactions
N_s	number of species in chemistry module
p_α	liquid/gas pressure, $[Pa]$
p_c	capillary pressure, $[Pa]$
p_b	entry pressure of capillary pressure model, $[Pa]$
P	pressure in thermodynamics, $[Pa]$
q_α	mass source term of phase α , $[kg/s]$
q_q^m	source term of basis species m in gas phase, $[mol/m^3]$
q_l^k	source term of basis species k in liquid phase, $[mol/kg\ w]$
Q	ion activity product
R	gas constant, $[J/K/mol]$
R_α	reaction term of phase α , $[kg/s]$

Table 1 Continued: Nomenclature

R_g^m	reaction term of basis species m in gas phase, $[mol/m^3]$
R_l^k	reaction term of basis species k in liquid phase, $[mol/kg\ w]$
S	concentration of species in chemical module
S_j	concentration of species j in chemical module
S_α	saturation of phase α
$S_{\alpha r}$	residual saturation of phase α
T	Temperature, $^\circ C$ and K in sec.
T_i	total concentration in chemistry module
\mathbf{u}_α	Darcy's velocity of phase α
v	molar volume, $[m^3/mol]$
V_α	volume of porous space occupied by phase α , $[m^3]$
V_{tot}	total volume, $[m^3]$
X_α^k	mass fraction of basis species k in phase α
\mathbf{x}	vector of primary variables of flow system
y_i	mole fraction of basis species i in gas phase
Z	compressibility factor

Greek Symbols	
$\alpha = \{l, g\}$	liquid/gas phase
α_{ij}	stoichiometric coefficient
β	power parameter of kinetic model
γ_k	activity coefficient
δ	power parameter of kinetic model
Δ	matrix of binary interaction coefficients of PR EOS
ε_g	gas quantity tolerance in reactive transport coupling
ε_{lin}	tolerance of GMRES
ε_{Nf}	residual function tolerance in flow coupling
ε_{qss}	quasi-stationary state tolerance in flow coupling
ε_{rt}	tolerance in reactive transport coupling
λ	parameter of Brooks-Corey model
μ_α	viscosity of phase α , $[Pa \cdot s]$
ρ_α	mass density of phase α , $[kg/m^3]$
ϕ_α	source velocity of phase α , $[m/s]$
τ_α	tortuosity of phase α
ϕ	porosity

Table 1 Continued: Nomenclature

φ_i^α	fugacity coefficient of species i in phase α
Ω	acentric factor set
\mathfrak{T}_α	transport operator of phase α
$\ \cdot\ _\infty$	infinity norm

Table 2: Abbreviations

AIM	adaptive implicit method
CFL	Courant-Friedrichs-Lewy number
DAE	differential algebraic equations based method
DSA	direct substitution approach
FIM	fully implicit method
FVM	finite volume method
GIA	global implicit approach
IMPEC	implicit pressure/explicit concentration
MMF	multiphase multicomponent flow
MMRF	multiphase multicomponent reactive flow
ODE	ordinary differential equations based method
OS	operator splitting
RT	reactive transport
SIA	sequential iterative approach
SNIA	sequential non-iterative approach

2. Governing processes and mathematical formulation

The entire isothermal MMRF problem is composed of phase α mass and component $c_{\alpha,k}$ mol conservation, mass balance, mass-action laws and other constitutive relations. The N_f fluid phases α , liquid and gas $N_f = 2$, are mobile, while solid phase is supposed to be immobile, $N_p = 3$. The chemical system of N_s chemical species and N_r linearly independent chemical reactions relies on Morel’s method [52] of N_c primary species that form a basis of all N_s species: $N_c = N_s - N_r$.

2.1. Mass conservation for each phase

We firstly introduce the standard multiphase compositional flow problem, the physical parameters and then present the alternative and its advantages.

Let us begin with the general mass conservation in terms of mass fraction $X_{\alpha,k}$ of species k in fluid phase α that forms $N_c N_f$ equations:

$$\frac{\partial(\phi S_\alpha \rho_\alpha X_{\alpha,k})}{\partial t} + \nabla \cdot (\rho_\alpha X_{\alpha,k} \mathbf{u}_\alpha - \rho_\alpha \mathbf{D}_{\alpha,k} \nabla X_{\alpha,k}) = R_{\alpha,k} + q_{\alpha,k}, \quad (1)$$

where ω is the porosity, S_α is the saturation of fluid phase α , ρ_α is the mass density of fluid phase α , \mathbf{u}_α is the Darcy-Muskat velocity of fluid phase α , $\mathbf{D}_{\alpha,k}$ is the dispersion tensor of fluid phase α , $R_{\alpha,k}$ is the reaction term and $q_{\alpha,k}$ is the external source term of species k in fluid phase α . The Darcy-Muskat law [53] states:

$$\mathbf{u}_\alpha = -\frac{k_{r\alpha}}{\mu_\alpha} \mathbb{K} (\nabla p_\alpha - \rho_\alpha \mathbf{g}), \quad (2)$$

where $k_{r\alpha}$ and μ_α are the relative permeability and the viscosity of fluid phase α , respectively, \mathbb{K} is the intrinsic permeability tensor, p_α is the pressure of fluid phase α , \mathbf{g} is the gravitational acceleration vector. The pressures are connected by $N_f - 1$ capillary pressure relations, the saturations and mass fractions sum to 1:

$$\sum_{\alpha} S_\alpha = 1, \quad (3)$$

$$\sum_k X_{\alpha,k} = 1, \quad (4)$$

that results in additional $2N_f$ constitutive relations. The conventional PDE system of isothermal compositional multiphase flow can be expressed from (1) by deriving a conservation of each species k in all phases \sum_{α} . These N_c nonlinear equations, $2N_f$ constitutive relations and $N_c(N_f - 1)$ phase equilibrium relations (sec. 2.3) give $2N_f + N_c N_f$ equations for $2N_f + N_c N_f$ unknowns $p_\alpha, S_\alpha, X_{\alpha,k}$. According to the Gibbs phase rule, the number of intensive properties is $N_c - N_p + 2$. So, if the number of phases is locally known, $N_p = 2$ for fixed temperature, at least N_c nonlinear equations (for pressure and composition) must be solved in order to establish the thermodynamic state. The geochemical system can be abundant in species that implies the solution of considerably large nonlinear systems.

In this work, we propose to handle the phase mass conservation system of N_f PDE whose size is independent of the number of chemical species N_s . We apply \sum_k to (1) to obtain

$$\frac{\partial(\phi S_\alpha \rho_\alpha)}{\partial t} + \nabla \cdot (\rho_\alpha \mathbf{u}_\alpha) = R_\alpha + q_\alpha, \quad (5)$$

taking into account that the sum of diffusive fluxes for each phase α is equal to 0. The external source term q_α can be also presented as

$$q_\alpha = \rho_\alpha \psi_\alpha, \quad (6)$$

where ψ_α is the source velocity. The system (5) of N_f equations, $N_f - 1$ capillary pressure relations and (3) are assembled for $2N_f$ unknowns p_α, S_α which are natural variables of multiphase flow problem.

When one of the phases disappears, the corresponding equation of Sys. (5) degenerates and the natural variables are inappropriate to describe the system. In this case, we pass to the single flow problem. Numerous formalisms are dedicated to the two-component two-phase flow and associated phase disappearance/appearance [1, 6, 10, 43, 65]. However, the liquid phase can be supposed to be present throughout the system, even if it remains in small amounts. Therefore, solving the system of N_f nonlinear equations (5) can be beneficial when $N_c > N_f$.

2.2. Mole conservation for each gas component and primary species

The liquid and gas phases consist of N_c primary species and N_g gas species, respectively, $N_g < N_c$, for which the transport must be solved. We chose the transport formulation in terms of concentrations $c_{\alpha,k} \propto \rho_\alpha X_{\alpha,k}/M^k$. Deriving the transport equations in mole/mass fractions as in Eq. (1), one neglects the density deviation $\nabla \rho_\alpha / \rho_\alpha$ that arises from the diffusive part of flux. Based on the primary species formalism [44, 75, 77, 89], the liquid transport is defined for total liquid mobile concentration $c_{l,k}$ of primary species k and the gas transport for gas concentration $c_{g,m}$ of gas species m , that evolves into $N_c + N_g$ transport (linear) equations:

$$\frac{\partial \phi S_l c_{l,k}}{\partial t} = \mathfrak{T}_l(c_{l,k}) + R_{l,k}(c_{l,k}, c_{s,k}) + q_{l,k}, \quad (7)$$

$$\frac{\partial \phi S_g c_{g,m}}{\partial t} = \mathfrak{T}_g(c_{g,m}) + R_{g,m}(c_{g,m}) + q_{g,m}, \quad (8)$$

where $c_{s,k}$ is the immobile concentration of species k , operator \mathfrak{T}_α includes the advective flux presented by Darcy-Muskat law (2) and, supposing non-Knudsen diffusion, Fick's law for diffusive-dispersive flux that yields [21, 44]:

$$\mathfrak{T}_\alpha(c_{\alpha,k}) = \nabla \cdot (\mathbf{D}_{\alpha,k} \nabla c_{\alpha,k} - c_{\alpha,k} \mathbf{u}_\alpha), \quad (9)$$

where $\mathbf{D}_{\alpha,k}$ involves the molecular diffusion $D_{\alpha,k}$, the tortuosity τ_α [49] and the longitudinal and transverse dispersion.

2.3. Mole balance for each primary basis species

Here, we introduce concisely the reactive part presented by CHESS [77], that yields the mass action law and phase equilibrium relations. As mentioned above, the reactive transport code is based on the primary species formulation, so we denote the concentration of species S_j , $j = 1, \dots, N_s$, that can be expressed as a function of basis species C_i , $i = 1, \dots, N_c$:

$$S_j \rightleftharpoons \sum_{i=1}^{N_c} \alpha_{ij} C_i, \quad (10)$$

where α_{ij} is the stoichiometric coefficient. At equilibrium, the mass action laws provide the reaction affinity:

$$S_j = \frac{K_j}{\gamma_j} \prod_{i=1}^{N_c} (\gamma_i C_i)^{\alpha_{ij}}, \quad (11)$$

where γ_j is the activity coefficient and K_j is the thermodynamic equilibrium constant of reaction j . All aqueous reactions are treated at equilibrium. For each basis species, the mole balances can be written by the total concentration T_i , $i = 1, \dots, N_c$, $T_i = \sum_{j=1}^{N_s} \alpha_{ji} S_j$,

$$T_i - C_i - \sum_{j=1, j \neq i}^{N_s} \alpha_{ji} \frac{K_j}{\gamma_j} \prod_{i=1}^{N_c} (\gamma_i C_i)^{\alpha_{ij}} = 0, \quad (12)$$

that constitute a system of N_c equations on C_i , $i = 1, \dots, N_c$.

2.3.1. Liquid mixtures

In the non-ideal liquid mixtures, the activity coefficients γ_i are not trivial and can be calculated by different models whose complexity increases with the concentration of solution, from less to more concentrated solution: truncated Davies formula [17], B-dot [32], SIT [30], Pitzer [61].

2.3.2. Gas-liquid equilibrium

The chemical potentials and also the corresponding fugacities of species i in mixture f_i^α are equal under equilibrium conditions. Considering the fugacity-activity ($\varphi - \gamma$) approach, it yields

$$Py_i \varphi_i^g = f_i = K_i^h \gamma_i x_i, \quad (13)$$

where y_i is the mole fractions of species i in gas, φ_i^g is the fugacity coefficient of species i , $K_i^h = K_i^h(T, P)$ is the corrected Henry's constant of species i , γ_i is the asymmetric activity coefficient of species i [48]. K_i^h involves the Poynting factor which corrects the reference fugacity in regard to the pressure and which is near the unity at low to moderate pressures. The fugacity coefficients can be calculated by cubic EOS, we use Peng-Robinson EOS [67] which precisely reflects the gas mixture properties for high pressure/temperatures. By given P , the cubic equation should be solved for the compressibility factor $Z = Pv/RT$, where v is the molar volume. The fugacity coefficient is then calculated [67], $\varphi_i^g = \varphi_i^g(T, P, Z, \mathbf{T}_c, \mathbf{P}_c, \mathbf{Z}_c, \mathbf{\Omega}, \mathbf{\Delta})$, where $\mathbf{T}_c, \mathbf{P}_c, \mathbf{Z}_c$ are sets of the critical temperature, critical pressure and compressibility factor of species in mixture, respectively, $\mathbf{\Omega}$ is the acentric factor set, $\mathbf{\Delta}$ is the matrix of empirical binary interaction parameters for each pair of species in mixture.

The EOS also provides the mass density for gas mixtures:

$$\rho_g = \frac{\sum_{i=1}^{N_g} y_i M_i}{v} = \frac{\bar{M}}{v}, \quad (14)$$

and by analogy for liquid mixtures with the $\bar{M} = \sum_{i=1}^{N_c} x_i M_i$.

It is worth mentioning that Raoult's law ($Py_i = P^{sat}x_i$) is derived from Eq. (13), supposing the low pressure condition and ideality of solution.

2.3.3. Liquid-solid equilibrium and kinetic relations

Considering that the activity of mineral equals unity and omitting index j , one derives from the mass action laws (11) the solubility product constant $K_{s,j}$ for solid at equilibrium

$$K_s = \prod_{i=1}^{N_c} (\gamma_i C_i)^{\alpha_{ij}} = 1/K. \quad (15)$$

However, to describe the non-equilibrium state of mineral, the ion activity product Q_s should be defined similarly to K_s but for the actual activities. Next, the saturation index SI yields the state of saturation by definition:

$$\text{SI} = \log \left(\frac{Q_s}{K_s} \right) = \begin{cases} < 0 & \text{undersaturated} \rightarrow \text{dissolution,} \\ = 0 & \text{saturated} \rightarrow \text{equilibrium,} \\ > 0 & \text{oversaturated} \rightarrow \text{precipitation.} \end{cases} \quad (16)$$

The kinetics of mineral precipitation/dissolution is usually modeled by the transition state theory rate laws [42]:

$$\frac{dS}{dt} = A_s k \left(\text{sign}(\text{SI}) \left(\left(\frac{Q_s}{K_s} \right)^x - 1 \right) \right)^y \prod_k a_k^{n_k} \quad (17)$$

where A_s is the specific surface area, k is the kinetic dissolution/precipitation rate constant, K_s is the equilibrium solubility constant for solid S , x, y, n_k are the fitting parameters, a_k is the activity of the potential catalyzing or inhibiting species.

The mole balance equations with the mass action laws for the species at equilibrium, the rate laws for kinetically limited solids and phase equilibrium relations constitute a complete system of algebro-differential equations whose solution yields the concentrations of basis and then secondary species. Other chemical reactions can also be handled by the formalism of basis species, e.g. cation exchange, surface and organic complexation.

3. Numerical solution

Applying OS concept in subsurface environmental modeling permits the independent development of separated parts of code and rigorous solution of each of them, that determines the majority choice of RT simulators [74]. The assessment of different coupling methods was studied on the MoMaS benchmark of RT codes [13, 12], during which the reliability both of SI and of GI approaches was revealed, the detailed results of HYTEC was demonstrated in [40]. Following the general strategy of integrating the (un)saturated flow in RT codes by OS, we solve the compressible two-phase flow block first, that involves the flow system, the gas transport equations and the EOS with the fluid properties models. Then the reactive transport coupling is applied. We propose to employ SIA for each of them, and hence there are, indeed, two internal SI couplings. Let us describe the applied methods for flow and transport discretization and the coupling methods afterward.

3.1. Discretization of flow and transport

The discretization of mass phase conservation (5) and mole species transport (7), (8) is built on a Voronoi-type finite volume method. The time approximation of two-phase flow (5) is fully implicit, the fluxes are handled

by TPFA. It was proven since the 1960s, that the transmissibility discretization affects the numerical stability and accuracy [4, 9, 72]; in this work, the interface coefficients of flow between adjacent cells are then evaluated implicitly and upstream. For the relative permeability $(k_{r\alpha})_{ij}$, the upstream space approximation is widely used, since its convergence was demonstrated on the Buckley-Leverett problem [7, 8]. A detailed comparison of temporal discretization presented in [7, 9] showed stability advantages of the implicit upstream treatment against explicit ones but increase of truncation errors at the same time. We will see the impact of truncation errors in Section 4.1. So, the relative permeability $k_{r\alpha}$, intrinsic permeability K , phase density ρ_α , phase viscosity μ_α for interface coefficient between the adjacent cells i, j at iteration $k + 1$ are defined as

$$(\cdot)_{ij}^{k+1} = \begin{cases} (\cdot)_i^k & \text{if } u_\alpha^k > 0, \\ (\cdot)_j^k & \text{else} \end{cases} \quad (18)$$

where $u_\alpha = \mathbf{u}_\alpha \mathbf{n}_{ij}$, \mathbf{n}_{ij} is the normal vector from i to j .

The discretization of density $(\rho_\alpha^g)_{ij}$ in the gravity term $\rho_\alpha \mathbf{g}$ needs to be treated differently: it is weighted with regard to the effective phase volume. If the gas phase is absent in one of the cells, the upstream treatment is applied [16]:

$$(\rho_\alpha^g)_{ij} = \begin{cases} \rho_{\alpha,i}\sigma + \rho_{\alpha,j}(1 - \sigma) & \text{if } (S_{\alpha,i} > 0) \wedge (S_{\alpha,j} > 0) : \\ & \sigma = V_{\alpha,i}/(V_{\alpha,i} + V_{\alpha,j}); \\ (\rho_\alpha)_{ij} & \text{else} \end{cases} \quad (19)$$

where $V_\alpha = \phi S_\alpha V_{tot}$, V_{tot} is the volume of the cell. The resulting nonlinear system is solved by Newton's method with an analytical Jacobian, that will be displayed below in Sec. 3.2.

The space discretization of transport operators (7), (8) is performed by the upstream weighting for advective flux and harmonic for effective diffusion, while one can apply the implicit or semi-implicit method in time. Applying a semi-implicit method for time discretization makes it possible to attain an accurate solution: implicit Euler scheme in time for diffusive-dispersive and Crank-Nicholson for advective part.

3.2. Coupling 1: Compressible two-phase flow

Since the phase flow system (5) is nonlinear, the classic Newton's method is applied. We denote the discretized equations (5) as $\mathbf{F}(\mathbf{x}) = 0$, where

$\mathbf{x} = (\mathbf{p}_l, \mathbf{S}_g)$. When the fluids are highly compressible (e.g. gas phase), the density properties should be precisely evaluated at each deviation of intensive variables, P, V, n , where n is the quantity of matter. The gas appearance and disappearance involve the density update in Newton's loop similar to the other flow parameters to ensure the implicit treatment of interface coefficients. However, the gas density can be strongly dependent on its composition. Therefore, the gas composition has to be calculated by employing the gas transport (8), denoted by $\mathbf{T}_g(\mathbf{c}_g; \mathbf{x}) = 0$. The flow coupling algorithm for time step $n + 1$ is presented in 1 and below its parsing.

Algorithm 1 Newton's method for Flow

```

1:  $\varepsilon_{Nf} = 1 \times 10^{-6}$ 
2:  $\varepsilon_{qss} = 1 \times 10^{-24}$ 
3:  $k_{max}^{fl} = 9$ 
4:  $k = 0$ 
5: while  $(\|\mathbf{F}(\mathbf{x}^k)\|_\infty \geq \varepsilon_{Nf} \|\mathbf{F}(\mathbf{x}^0)\|_\infty) \wedge (\|\mathbf{F}(\mathbf{x}^k)\|_\infty \geq \varepsilon_{qss})$ 
    $\wedge (k \leq k_{max}^{fl})$  do
6:   find  $\delta \mathbf{x}^{k+1}$ :  $\mathbf{J}(\mathbf{x}^k) \delta \mathbf{x}^{k+1} = -\mathbf{F}(\mathbf{x}^k)$ 
7:    $\mathbf{x}^{k+1} \leftarrow \mathbf{x}^k + \delta \mathbf{x}^{k+1}$ 
8:   find  $\mathbf{c}_g^{k+1}$ :  $\mathbf{T}_g(\mathbf{c}_g^{k+1}; \mathbf{x}^{k+1}) = 0$ 
9:   update EOS and physical parameters
10:   $k \leftarrow k + 1$ 
11: end while

```

The user defined or default tolerance $\varepsilon_{Nf} \in [10^{-8}, 10^{-6}]$ and maximum number of Newton iteration k_{max}^{fl} for the flow coupling are firstly initialized. Next, the linearized system is solved for the increment $\delta \mathbf{x}$, line 6. All the partial derivatives involved in the Jacobian \mathbf{J} of discretized flow system are analytical. For ex. the derivative of gas density interface coefficient is expressed by

$$\frac{\partial(\rho_g)_{ij}}{\partial p_{g,i}} = \frac{\partial(\rho_g)_{ij}}{\partial \rho_{g,i}} \frac{\partial \rho_{g,i}}{\partial p_{g,i}}, \quad \frac{\partial \rho_{g,i}}{\partial p_{g,i}} = -\frac{\bar{M}_i}{v_i^2 (\partial p_g / \partial v)_i}, \quad (20)$$

where $\partial p_g / \partial v$ is the analytical derivative arising from the corresponding EOS. One can note that the density derivatives are composition dependent and proportional to the average of molecular weight \bar{M} as well as the density function Eq. (14). Various solvers exist for solving multiphase system of

linear equations [14], we apply GMRES [68], that is one of the most prevalent and efficient methods, and ILU(0) [79] as a preconditioner.

With the velocities and saturations given by \mathbf{x}^{k+1} from step 7, the linear transport system $\mathbf{T}_g(\mathbf{c}_g^{k+1}) = 0$ is solved for \mathbf{c}_g^{k+1} at line 8 by using GMRES and ILU(0). Due to the modified composition, the EOS parameters must be updated in order to evaluate a new molar volume v by solving the EOS analytically. Then, the physical properties can be calculated, line 9. Thereafter three stop criteria must be checked, line 5: two for the flow system residual and one for the number of iterations.

The proposed coupling 1 corresponds to Newton's family for the flow system in regard to \mathbf{x}^{k+1} , while it can be seen as Picard's method (fixed point method) for the transport equations on \mathbf{c}_g^{k+1} . Inasmuch as the gas phase is supposed to be compressible and the significant difference in gas density may occur over all modeled domain, one can include an additional criterion for gas quantity n_g deviation:

$$\max \frac{|n_g^{k+1} - n_g^k|}{n_g^{k+1}} \leq \varepsilon_g, \quad (21)$$

where max is maximum over the modeled domain. After numerous tests, we deduce, that this can be not necessary condition but sufficient to finish the loop that depends on the complexity of gas dynamics. In spite of neglecting the criterion (21), there is no lack of solution accuracy. In addition, the reactive transport coupling 3.3, that goes after the flow coupling 1, entails the convergence (stop) conditions for gas and solid phase and guarantee the conservation of the whole system.

An adaptive time-stepping is implemented in regard to the relaxed CFL condition, number of Newton iterations and maximum saturation/pressure changes. Extremely large time steps are not reasonable, even if the scheme is unconditionally stable. Moreover, the following reactive transport can require smaller time step. In this case, the smaller inner time stepping is usually used. Nevertheless, when the RT coupling is finished, the aggressive chemistry would provide the important changes returning the larger reaction terms and the stronger modified fluid and rock properties to the flow block at the next time step. So, the flow solution would be further from the first guess and consequently, the convergence rate would decrease as it is quadratic only near the root. This subject will be discussed in Sec. 3.4.1.

Notice that there is no calculation of geochemical reactions in this coupling, but the reaction term R_α , that can be simply expressed by:

$$R_{\alpha}^n = \frac{(V_{\alpha}\rho_{\alpha})^{n,rt} - (V_{\alpha}\rho_{\alpha})^{n,fl}}{t^{n+1}}, \quad (22)$$

where t^n is time step n , $(\cdot)^{n,fr}$ and $(\cdot)^{n,rt}$ denote the values obtained by the flow and reactive transport couplings, respectively. So, it is then estimated *a posteriori* at each time step and remains the mass conservation throughout the calculations. It is also worth mentioning, that when three phases, g, l, s , are in the system, the term R_g reflects not only the gas and liquid connection but also the implicit gas-liquid-solid mass transfer, which is not negligible in conventional geochemical modeling. Precipitation and dissolution of some minerals can lead to gas depletion or formation, that impacts on saturation and pressure. These phenomena were numerically demonstrated in [29]. Consequently, the value of reaction term can be of great importance, depending on the physical and chemical problem statement and on the aimed accuracy. *E.g.* in Vostrikov’s work, the reaction terms between the flow for dominant components and the reactive transport were neglected, given the next assumptions: “only very small amounts of minerals are transferred to the liquid form” and “minor components do not have a significant impact on the physical parameters of system”.

Algorithm 1 constitutes simple solving the two-phase system, where the gas concentrations are assumed to be the secondary variables just like the relative permeability, the capillary pressure and others. In this coupling, the flow can be fully unreactive supposing that the gas displacement is the dominant mechanism and the phase equilibrium calculations are taken explicitly from the reactive transport coupling, in this case.

Pursuing the idea of solving the phase flow formulation (5) and not the compositional problem, other options of coupling were tested, for example: Step 1. the phase conservation system to trace the first estimation of velocities and saturations; Step 2. the reactive transport coupling and finally Step 3. the phase flow problem, solving Newton’s method for the second time, but taking into account the phase transfer due to the geochemical reactions. This method demonstrated disadvantageous effects on the convergence and performance because of the delayed treatment of the fluid properties at Step 1. Thus, the gas properties need to be implicitly estimated in the iterative loop of flow to guarantee the convergence and unconditional stability, especially if in the aimed application, the mechanical displacement of gas front occurs, so the advective dominant flow takes place. As a result, the flow system is fully implicit and inherits its advantages in contrast to the time-step restriction

of IMPES method.

3.3. Coupling 2: Reactive transport

Relying on Algorithm 1 allows to find the phase velocities and saturations and consequently, to manage the reactive transport problem which consists of the linear transport and nonlinear chemical equations. So, analogous to the standard coupling in saturated porous media, Picard's method can be employed. On the one hand, the gas and liquid transports can be summed and therefore written for each species similar to GIA for multiphase multi-component flow [15, 43, 44] by applying phase appearance and disappearance criteria and K-values $K_i = y_i/x_i$, that would be $K_i = K_i^h \gamma_i / (P \varphi_i^g)$ in this work, derived from (13). In the non-ideal approach, γ_i is calculated regarding electrostatic state of ions through the ionic strength, furthermore, φ_i^g is the function of the whole mixture properties including y_i itself, then K_i depends both on aqueous and on gaseous composition and should be updated at each iteration as secondary variable. On the other hand, remaining liquid (7) and gas (8) operators apart, one can apply the gas concentration from Algorithm 1 as a first guess and avoid K-value calculation by introducing the reaction terms. By denoting transport operators by $\mathbf{T}_\alpha(\mathbf{c}_\alpha) = 0$ and geochemical reactions, described in Sec. 2.3 by $\mathbf{R}(\mathbf{c})$, where $\mathbf{c} = \{\mathbf{c}_l, \mathbf{c}_g, \mathbf{c}_s\}$, we propose the SI reactive transport algorithm 2.

Algorithm 2 Picard's method for Reactive Transport

```

1:  $\varepsilon_{rt} = 1 \times 10^{-5}$ 
2:  $k_{max}^{rt} = 60$ 
3:  $k = -1$ 
4: while  $\left[ \left( \max \frac{|c_g^{2k+2} - c_g^{2k}|}{c_g^{2k+2}} \geq \varepsilon_{rt} \right) \vee \left( \max \frac{|c_s^{2k+2} - c_s^{2k}|}{c_s^{2k+2}} \geq \varepsilon_{rt} \right) \right] \wedge (k \leq k_{max}^{rt})$ 
   do
5:    $k \leftarrow k + 1$ 
6:    $\mathbf{T}_g(\mathbf{c}_g^{2k+1}) = 0$ 
7:    $\mathbf{T}_l(\mathbf{c}_l^{2k+1}) = 0$ 
8:    $\mathbf{c}^{2k+2} = \mathbf{R}(\mathbf{c}^{2k+1})$ 
9: end while

```

Let us set out the specific aspects of the SI reactive transport algorithm 2. The reaction term, devoted to replicating dissolution and evaporation rate,

is introduced in gas transport operator at line 6. Omitting the porosity and the external sources, it can be formulated for gas species i as

$$\frac{S_g^{n+1}c_{g,i}^{n+1,2k+1} - S_g^nc_{g,i}^n}{t^{n+1}} = (\mathfrak{T}_{g,i})_h + \frac{S_g^{n+1}}{t^{n+1}}r_{gl,i}^{n+1,2k+1}, \quad (23)$$

$$r_{gl,i}^{n+1,2k+1} = d_i^{n+1,2k}c_{g,i}^{n+1,2k+1} + e_i^{n+1,2k}, \quad (24)$$

where $(\cdot)_h$ denotes the discretization, $d_i^{n+1,2k}$ and $e_i^{n+1,2k}$ reflect the dissolution and evaporation processes, respectively. The terms $d_i^{n+1,2k}$ and $e_i^{n+1,2k}$ are calculated at the previous iteration $2k$, while $r_{gl,i}^{n+1,2k+1}$ corresponds to the iteration $2k+1$. Considering the concentrations obtained on the previous iterations $\{1, \dots, 2k\}$, these parameters are defined as

$$d_i^{n+1,2k} = \sum_{m=1}^k \mathbb{1}_{\mathbb{R}_{>0}}(\Delta c_{g,i}^{n+1,2m}) \frac{\Delta c_{g,i}^{n+1,2m}}{c_{g,i}^{n+1,2m}}, \quad (25)$$

$$e_i^{n+1,2k} = \sum_{m=1}^k \mathbb{1}_{\mathbb{R}_{\leq 0}}(\Delta c_{g,i}^{n+1,2m}) \Delta c_{g,i}^{n+1,2m}, \quad (26)$$

where

$$\Delta c_{g,i}^{n+1,2m} = c_{g,i}^{n+1,2m} - c_{g,i}^{n+1,2m-1} \quad (27)$$

and $\mathbb{1}_{\mathbb{R}_{>0}}(\cdot)$ is the indicator function of the set of strictly positive real numbers. Note that equation (28) is still linear and can be directly solved by one of the linear solvers, GMRES with ILU(0) for example. This new distribution of gas species i entails different mass transfer that should be associated with the corresponding total liquid mobile concentration of primary species j . Then, the liquid transport operator, line 7, is defined by

$$\frac{S_l^{n+1}c_{l,j}^{n+1,2k+1} - S_l^nc_{l,j}^n}{t^{n+1}} = (\mathfrak{T}_{l,j})_h + \frac{S_l^{n+1}}{t^{n+1}}r_{lg,j}^{n+1,2k+1} + R_{ls,j}(c_{s,j}), \quad (28)$$

$$r_{lg,j}^{n+1,2k+1} = \sum_i \alpha_{ij}r_{gl,i}^{n+1,2k+1}, \quad (29)$$

where $R_{ls,j}$ is the reaction term of liquid-solid interaction whose detailed description and variable porosity management can be found in [41].

Following the transport, the reactive part aims at finding a new local equilibrium state, line 8. The nonlinear chemical system for basis species is solved at each cell by Newton's method with the line search procedure. The

Jacobian is analytically evaluated. To improve the convergence, a new basis can be chosen independently in each cell, hence its set of primary equations also changes. Then, the gas and solid concentrations changes are subjected to stop criterion verification, line 4, if the phase interactions, liquid-gas, liquid-solid, take place.

3.4. Coupling between compressible two-phase flow and reactive transport

3.4.1. Type of coupling

The sequential coupling of the flow 1 and reactive transport 2 modules has a specific limitation arising from the former, which is supposed unreactive. The flow method was initially devised for modeling the gas appearance due to the mechanical displacement but not due to the phase transition, from liquid to two-phase state. Nevertheless, within the reactive transport module, the evaporation, dissolution and other chemical processes are permitted and consequently, it changes the occupied volume and mass of each phase. Also, the gas phase can vanish or appear. The impact of such modifications varies with the chemical system of problem. We manage the changes arising from the RT in the flow block *a posteriori*, by updating the fluid and rock properties and the explicit reaction terms R_α , Eq. 22. This approach is valid if chemical reactions rate and time steps remain small; otherwise, the severe changes in the fluid and rock properties hinder convergence of Newton's method, as initial guess is far from the solution. Moreover, the explicit treatment of geochemical impact on the flow can be significantly underestimated. To overcome this problem and to model the reaction driven advection, the tight coupling, similar to that of STOMP[85], should be applied in the case of the high change rate. It permits to update the velocities and the fluid/rock properties implicitly: iterative procedure of flow and SI reactive transport. However, when the mechanical force is dominant, it is computationally efficient to use the sequential coupling between the flow and the RT module.

Applying GIA avoids this issue by definition, since the equilibrium state is calculated at each Newton iteration by using the thermodynamics flash (Rachford-Rice equation) [48] or the complementary conditions. Instead, the compositional flow system consists of a larger set of nonlinear equations than that of the proposed method. Furthermore, in case of variable switching, the primary variables should be adapted to the local equilibrium state, and it requires to rebuild the Jacobian matrix.

3.4.2. Numerical assessment

By means of this approach the whole problem is divided into two subsystems: the compressible flow coupling 1 composed by N_f nonlinear equations of flow and N_g linear equations of gas transport, then the reactive transport coupling 2 of $N_c + N_g$ linear transport equations and $N_c + N_{kin}$ nonlinear equations from chemical system. Let us estimate the computational impact of each part. Supposing the uniform time step both for flow and for reactive transport, we denote the calculation time of the whole system t , the flow coupling per iteration t_{flc} , the flow operator t_{fl} , the gas transport t_{gt} and for the reactive transport per iteration t_{rtc} , then

$$t = N_{Nit}t_{flc} + N_{Pit}t_{rtc} = N_{Nit}(t_{fl} + t_{gt}) + N_{Pit}t_{rtc}, \quad (30)$$

where N_{Nit} is the number of Newton iterations, N_{Pit} is the number of Picard iterations. By taking into account $t_{gt} \ll t_{fl}$ and $N_{Pit}t_{rtc} \in (t_{fl}/10^3, N_{Nit}t_{fl}]$, we deduce that the calculation time per time step for this work can be expressed by following

$$N_{Nit}t_{fl} < N_{Nit}t_{fl} + N_{Pit}t_{rtc} \leq 2N_{Nit}t_{fl}, \quad (31)$$

where the lower border corresponds to the problem with a low geochemical complexity (*e.g.* CO_2 and H_2O) and the calculation time t is mostly based on the flow operator part, making t_{fl} decisive factor, as a result. It can be reduced by performant linear solver and preconditioner [37] or/and by using the AIM for the flow system. The upper limit can be reached, when the aggressive geochemistry is modeled with the dis-/appearance of solids, gases, the redox reactions etc. Although, the calculation time t is still strongly dependent on t_{fl} . Thus, we conclude that the more the chemical system is abundant in components, the more the proposed flow coupling is advantageous in comparison with the methods based on the component flow formulation, considering that the larger time stepping in GIA can be compromised by the required reactive time step.

The inner couplings have their own tolerance, another benefit of the OSA: $[10^{-8}, 10^{-6}]$ for flow, 10^{-5} for reactive transport and 10^{-12} for the chemistry, that results in the mass balance error in the range $[10^{-6}, 10^{-5}]$ according to the test cases carried out for different types of geochemical and hydrodynamic complexity.

4. Numerical simulations

The proposed method is first tested by modeling a problem with the self-similar solution. Next, we apply it to the simulation of CO_2 injection that permits to evaluate the capabilities of the simulator to represent the physical behavior and its computational efficiency.

4.1. 1D axisymmetric problem: radial flow from a CO_2 injection well

The axisymmetric problem of constant injection in a saturated, horizontal, infinite reservoir admits the self-similar variable R^2/t , whose property makes it an ideal way to validate the numerical code. In addition, this advantage was availed to model a constant 100 kg/s CO_2 injection in a long aquifer $100000 \times 100 \text{ m}^2$, in the workshop “Intercomparison of numerical simulation codes for geologic disposal of CO_2 ” initiated by Lawrence Berkeley National Laboratory [64]. We use the parameters presented in [64], adapting some of them: as the pore compressibility is neglected, we intensify the intrinsic permeability, $K = 2 \times 10^{-13} [\text{m}^2]$; the water disappearance is prohibited; the fluid properties and the solubility are treated differently. In HYTEC, the gas and liquid densities are provided by the Peng-Robinson models [2, 36, 67], the viscosities are predicted by [5, 34]. The partial molar volume at the in-

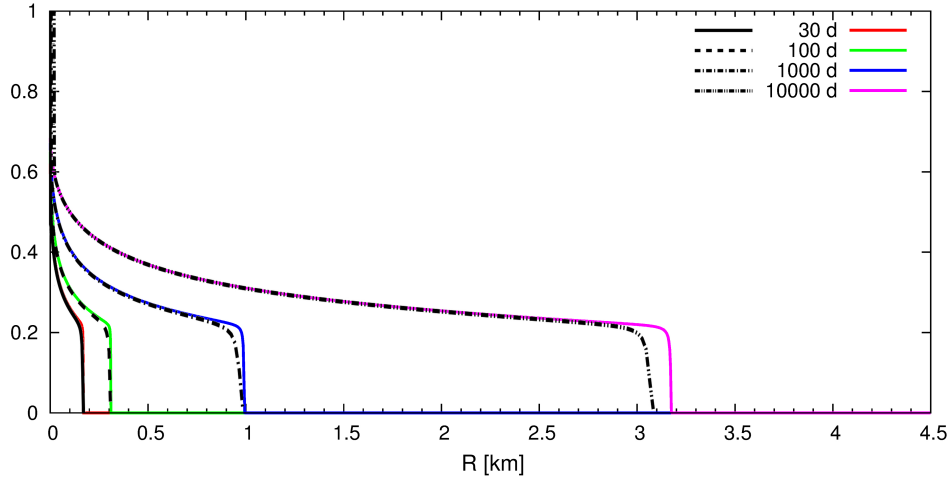


Figure 1: 1D axisymmetric problem: gas saturation S_g . Results of HYTEC (in color) and of TOUGH2-ECO2, [64] (black)

finite dilution in the water is averaged over the relevant pressure range; the aqueous activities and gaseous fugacities are simulated according to the b-dot and the PR78 [67] models, respectively. In [64], TOUGH2-ECO2 module was used, whose description is given in [63].

Despite the mentioned deviation in parameters, the gas saturation front is similar in both cases, especially over the first 1000 d, Fig. 1. The increased discrepancy in saturation front position at 10000 d arises from the water disappearance zone modeled in [64]. Notice that the the saturation curves of HYTEC are almost perpendicular to the R-axis in contrast to those of TOUGH2-ECO2, their slope grows with time that is just the truncation error effect. The results of gas saturation as a function of R^2/t are illustrated in Fig. 2 and demonstrate a high accuracy of the numerical code.

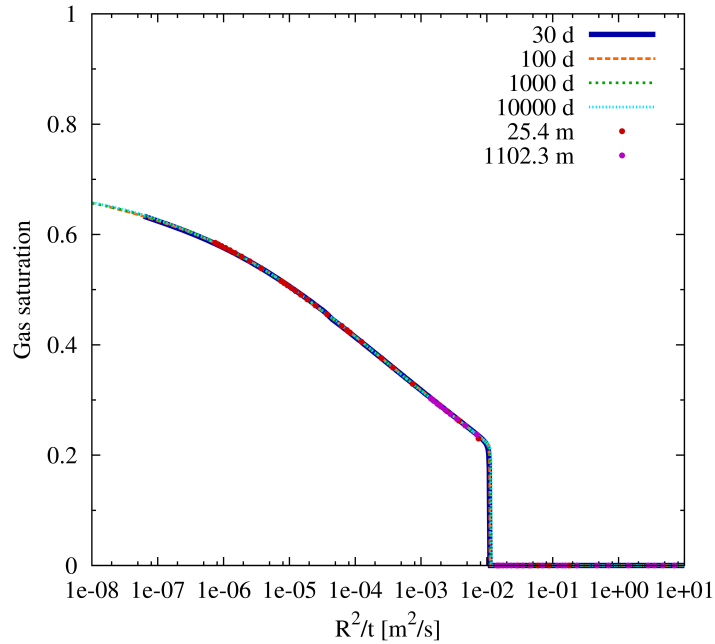


Figure 2: 1D axisymmetric problem: gas saturation S_g as a function of R^2/t

4.2. 2D problem: CO_2 injection in a fully water-saturated domain

The next 2D problem of CO_2 injection in a fully water-saturated domain was proposed by Neumann et al. (2013). The injection of constant rate

0.04 kg/m²/s is modeled through the left bottom boundary (whose length was not provided in [55]) of the rectangle reservoir 600×100 m². In this work, the injection border is 10 m long, then the debit is 0.4 kg/s. The Dirichlet boundary conditions are set at the right border of the reservoir: hydrostatic pressure and $S_g = 0$. By neglecting the dispersion and assuming the Millington-Quirk tortuosity model [49], the effective diffusion for this problem takes the following form:

$$D_\alpha^e = \omega^{4/3} S_\alpha^{10/3} D_\alpha. \quad (32)$$

The geometry and grid dimension are taken after Neumann et al., while the fluid properties and the solubility differ from the reference. The modeling of fluid properties and of phase equilibria is analogous to that given in Sec. 4.1.

The fluid dynamics is represented exactly as in the reference [55]: the CO₂ forms a bubble that grows and rises upward, then the current is distributed

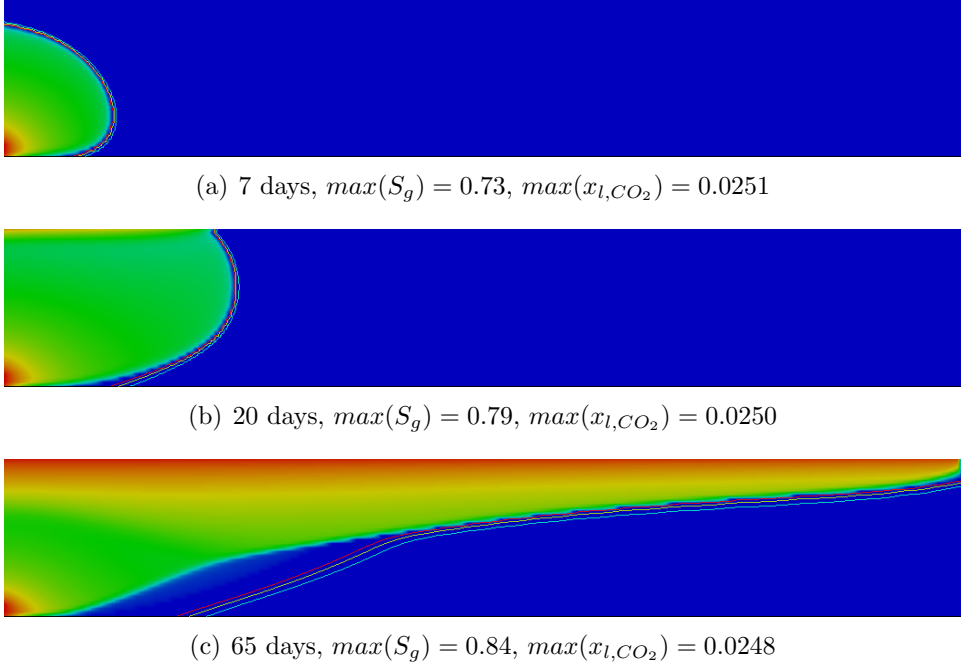


Figure 3: 2D problem of CO₂ injection in a fully water-saturated domain: gas saturation from $[0, \max(S_g)]$ and contours of CO₂ mole fraction x_{l,CO_2} at 0.005, 0.011, 0.016

along the top of the aquifer by gradually developing its area, Fig. 3. The gas saturation is lower than that of the reference performed by DUNE, the mole fraction of CO_2 x_{l,CO_2} is higher. This might be due to an uncertainty on the injection rate used in [55] and due to the different fluid properties and phase equilibria models.

The same grid dimension as in [55] is chosen for this simulation, 240×40 cells. The maximum number of Newton iterations is set to 9 in HYTEC for this problem. The tolerance of Newton’s method (Algorithm 1) is $\varepsilon_{Nf} = 1 \times 10^{-7}$ and the gas quantity criterion Eq. (21) is $\varepsilon_g = 1 \times 10^{-5}$. The mass conservation error is of order 10^{-6} .

The initial time step is set to 156 s, which is also the minimum time step $\min(dt)$. The time stepping for HYTEC and DUNE is presented in Table 3. The HYTEC’s imposed maximum time step $\max(dt)$ is taken slightly higher (by 8%) that results in a larger average dt and in a faster execution of HYTEC, by at least 9%.

In Table 4, the execution time of HYTEC includes the grid construction, initialization, output printing, solvers, secondary property modules. During our simulation, the total number of time steps (successful and unsuccessful) is 39% less than that of DUNE, that can be explained by slightly higher average time step and rate of Newton’s method convergence in HYTEC.

By employing our method, the unsuccessful steps appear right after the moment when the gas current reaches the right boundary, hence it is restricted to the further $\text{CO}_2(\text{g})$ propagation. Nevertheless, the dissolved CO_2 can be released; the depressurization thus increases the physical complexity of the system, hence the unsuccessful Newton iterations.

Table 3: 2D problem of CO_2 injection in a fully water-saturated domain: time stepping

	$\min(dt)$, s	$\max(dt)$, s	$\text{mean}(dt)$, s
DUNE	156.25	5000	3579.7
HYTEC	156	5400	4474.9

Table 4: 2D problem of CO₂ injection in a fully water-saturated domain: information of Newton successful and unsuccessful iteration number (Ni), total number of steps (successful and unsuccessful) and total execution time

	Tot. of time steps	$mean(Ni)$	Tot. of Ni	Tot. exec. time, s
DUNE	2249	3.9	-	13975
HYTEC	1380	5.2	7183	12708

5. Conclusion

A new solution method of compressible multiphase flow was proposed, that can be integrated as a module in conventional reactive transport frameworks, based on the operator splitting or global implicit approach. The flow method is composed by the phase conservation formulation, the gas transport and the equations of state. The versatile structure allows to conserve the constant number of nonlinear equations of the flow problem, independent of the modeled geochemical system, while chemistry basis can change during one time step to enhances the convergence. This characteristic makes it advantageous for modeling the multicomponent problems. The flow phase system can be also presented by other formulations of primary variables, *e.g.* pressure/capillary pressure. It is worth mentioning that it is possible to generalize the variable switching in rigorous way. The whole flow coupling preserves the fully implicit advantages of the multiphase flow discretization. The present method was implemented in the SIA based reactive transport simulator HYTEC. The numerical code was verified by modeling the benchmark problem admitting the self-similar solution, the computational efficiency was showed by the simulation of CO₂ injection and compared with that of DUNE.

Acknowledgements

This work was supported by MINES ParisTech and BRGM. I. Sin thanks Christophe Coquelet (Centre Thermodynamique des Procds, MINES ParisTech) for his perspicacious suggestions on the thermodynamics development and Claude Tadonki (Centre de Recherche en Informatique, MINES ParisTech) for the constructive discussions on the linear solvers.

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3.3 Verification

Once the numerical code is built, it should pass the prerequisite verification steps:

- mass balance conservation,
- modeling an analytical solution,
- benchmarking, reference models.

In Sec. 3.2, we have demonstrated the code capabilities to represent the semi-analytical and reference models. Here, we give additional information on the numerical results presented above, Sec. 3.3.1 and 3.3.2. Then we propose a benchmark study, Sec. 3.3.3. The mass conservation was calculated for all modeled problems in this work and the order of relative error is less than 10^{-6} .

3.3.1 1D axisymmetric problem: radial flow from a CO₂ injection well, Pruess et al. (2002)

The parameters of the model can be found in Table 2.1. The fluid densities are given by the PR78 and T-VTPR models (see Eq. (4.1.19) and Sec. 4.2) for which the binary interaction parameter are calculated by the PPR78. The gas viscosity is predicted by Altunin75, Eq. (4.3.8); we use the Islam-Carlson model of the liquid viscosity, Eq. (4.3.12). Modeling the 1D radial problem with self-similar solution by the full coupling reveals the growing impact of solubility and compressibility, compare with Figs. 2.4(a), 2.5 and 2.6.

3.3.2 2D problem: CO₂ injection in a fully water-saturated domain, Neumann et al. (2013)

Neumann et al. (2013) posed the model of CO₂ injection in homogeneous media taking the gravity into account. The geometry and grid dimension are the same as those of Neumann et al., while the fluid properties and the solubility differ from the reference. Table 3.1 contains the used model parameters. The tortuosity is presented by the Millington-Quirk model. The density and viscosity are variable and their models were presented in Sec. 4.3. During 65 days, 2.25 kt of CO₂ was injected.

Table 3.1 – Parameters of the 2D problem of CO₂ injection (Neumann et al., 2013).

Initial Conditions		
$\Omega = [0, 600] \times [0, 100] \text{ [m}^2\text{]}$		
$\Omega :$	$p_{l0} = p_{atm} + \rho_l g (900 - z) \text{ [Pa]},$	$S_{g0} = 0$
Temperature	40 °C	
Boundary Conditions		
$\Gamma^N = [(0, 0); (0, 10)] \text{ [m]}$		
$\Gamma^D = [(600, 0); (600, 100)] \text{ [m]}$		
$\Gamma^N :$	$\psi_l = 0,$	$\psi_g = 4 \times 10^{-2} \text{ [kg/m}^2\text{/s]}$
$\Gamma^D :$	$p_{l0} = p_{atm} + \rho_l g (900 - z) \text{ [Pa]},$	$S_g = 0$
$\partial\Omega \backslash (\Gamma^D \cup \Gamma^N):$	$\psi_l = 0,$	$\psi_g = 0$
Matrix Properties		
Porosity ϕ	0.2	
Permeability K	$10^{-12} \text{ [m}^2\text{]}$	
Fluid-Matrix Properties		
Brooks-Corey capillary pressure and relative permeability		
Eq. 2.1.7, (2.1.9) and (2.1.10): $p_b = 10^3 \text{ Pa}, \lambda = 2, S_{lr} = 0, S_{gr} = 0$		
Fluid Properties		
Density	ρ_l : T-VTPR	ρ_g : PR78
Viscosity	μ_l : Islam-Carlson	μ_g : Altunin75
Diffusion coefficient	$D_l = D_g = 2 \times 10^{-9} \text{ [m}^2\text{/s]}$	

3.3.3 Benchmark proposition

In the previous sections, we implemented our coupling method for MMF and RT (3.2). Some validation work was performed either based on analytical solutions or literature results. A full description of the thermodynamic laws was then presented, including some applications to the $\text{CH}_4\text{-CO}_2\text{-H}_2\text{S-H}_2\text{O}$ system (4.2).

Based on these developments, a new application is proposed. The application is built from the work by Bonnaud (2012); Bonnaud et al. (2012): the authors attempt to simulate a mechanism possibly responsible for the creation of heterogeneities in the composition of gas reservoirs. In their scenario, an initially sour reservoir is leached by an underlying aquifer; the differential solubility of the gas components leads to local modifications of the gas composition. The system includes a tight connection between transport and geochemistry. The evolution of the mass/volume due to the reactive transport affects the pressure field, which leads to a strong feedback of chemistry on flow. An accurate modeling of a multicomponent gas phase is also required for this problem. Finally, the complexity of the CO_2 and H_2S geochemistry in the host rock is of particular interest.

Result analysis based on physical meaning also reach its limits due to the numerous couplings and feedbacks. We therefore decided to use this problem as the basis for a numerical benchmarking exercise, open for the reactive transport modeling community. The full description of the context and the increasingly complex exercises proposed in the benchmark are detailed in the following paper Sin et al. (submitted), provided below. The numerical results are partly reported in Sin et al. (2015).

2D simulation of natural gas reservoir by multiphase multicomponent reactive flow and transport – description of a benchmarking exercise

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Abstract

Different ways of coupling are used to handle the mathematical and numerical difficulties of multiphase multicomponent flow and transport coupled with geochemistry. In absence of analytical solutions for such complex systems, the benchmarking of reactive multiphase simulators is essential for verification and validation of existing methods. This paper presents a reactive transport benchmarking exercise which aims at specifically targeting the numerical capacity of the codes in the context of liquid and gas flow, multicomponent liquid and gas transport, gas solubility, and gas-water-rock interaction. The exercise is based on a description of the long term evolution of the sour gas reservoir. The set of benchmark cases were established to study geochemical, thermo- and hydrodynamical mechanisms.

Keywords: multiphase flow, acid gas leaching, natural gas reservoir, gas mixtures, numerical modeling, benchmarking

1. Introduction

Reactive transport codes are quantitative tools that allow to explore the behavior of complex geoscience systems: coupling and feedbacks between flow

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and transport of fluids in the pore space and reactions between aqueous species,
 5 mineral surfaces, rocks and a possible gas phase. They rely on the resolution
 of several sets of equations describing the hydrodynamics and the geochem-
 istry of the system [1]. Reactive transport simulation are now used in a wide
 variety of applications, in numerous domains and at different scales. *E.g.*, the
 HYTEC code, developed at MINES ParisTech [2, 3], has been applied to cement
 10 degradation [4], radioactive waste disposal (performance assessment, near- and
 far-field processes [5, 6]), geological storage of acid gases [7, 8, 9], uranium *in*
situ recovery processes [10].

For the gas-water-rock interaction (e.g. for the simulation of enhanced gas
 or oil recovery, gas storage, CO₂ sequestration), the models are based on cou-
 15 pled, non-linear equations for multiphase multicomponent reactive flow and
 transport coupled with chemistry and water-gas equilibrium. Different ways
 of coupling are used to handle the mathematical and numerical difficulties
 [11, 12, 13, 3, 14, 15]. The significant mathematical and computational chal-
 lenges impose constant verification of the accuracy of the codes [16]. Unfor-
 20 tunately, due to the non-linearity and numerous couplings, analytical solutions
 are limited to very specific and overly simplified or de-coupled test cases [e.g
 2, 17]. Alternatively, validation on experimental data (from laboratory or field)
 focuses on the accuracy of the physical and chemical laws included in the codes,
 rather than the accuracy of the numerical schemes. Inter-comparison between
 25 codes is therefore a key for code verification.

Several such benchmarking exercises have been promoted in the recent lit-
 erature [18, 16, 19, 2, 20]. Also noteworthy is the subsurface environmental
 simulations benchmarking initiative [21, 22], which aims at promoting an inter-
 national community around reactive transport benchmark exercises definition
 30 and resolution.

This paper presents such an reactive transport benchmarking exercise. The
 aim is to specifically target the numerical capacity of the codes in the context
 of multiphase (water and gas) flow, with several coupling between water and
 gas transport, multicomponent gas solubility, and gas-water-rock interaction.

35 The exercise stands half-way between pure numerics and direct application: the overall description should be geologically realistic, the multiple couplings should be preserved so that the verification is demonstrative, and finally the complexity should be moderate in order to keep the computational intensity as reasonable as possible.

40 The exercise is based on the description of the long term evolution of a sour gas reservoir. The geometry was simplified and several simplifying hypotheses are taken to allow for different complexity applications. A general overview of the geological context is proposed in the following section: the focus is on the physical and geochemical processes at stake in the system. Then, 45 the usually-accepted equations describing these processes are presented, with a view to identifying the mathematical bases of the system and highlighting the couplings. A description of the exercises is then given: the actual geometry, parameters, initial and boundary conditions are detailed. Finally, some insights to the results (using the HYTEC code) are given: however, the paper does 50 not provide a comprehensive description of the results to allow some liberty for potential benchmarks.

Table 1: Nomenclature

Latin Symbols	
a	attraction parameter in Peng-Robinson EOS
a_α	dispersivity in phase α , $[m]$
A_s	specific surface area, $[m^2/m^3 \text{ solution}]$ or $[m^2/kg \text{ mineral}]$
b	van der Waals covolume
c_i^l	total liquid mobile concentration of basis species i , $[mol/kg \text{ w}]$
\bar{c}_i	immobile concentration of basis species i , $[mol/kg \text{ w}]$
c_i^g	gas concentration of basis species i , $[mol/m^3]$
C_i	concentration of primary species i in chemical module
D_α	molecular diffusion coefficient of phase α , $[m^2/s]$
D_α^{eff}	effective diffusion coefficient of phase α , $[m^2/s]$

\mathbf{g}	gravitational acceleration vector, $[m/s^2]$
k	kinetic constant, $[mol/m^2/s]$
$k_{r\alpha}$	relative permeability of phase α
$\mathbb{K} = KI$	intrinsic permeability tensor, $[m^2]$
K_j	solubility constant
K_s	solubility constant of solid phase
M	molecular weight, $[kg/mol]$
N_p^{tr}	number of basis species in transport module
N_p	number of basis species in chemistry module
N_s	number of species in chemistry module
p_α	liquid/gas pressure, $[Pa]$
p_c	capillary pressure, $[Pa]$
p_e	entry pressure of capillary pressure model, $[Pa]$
P	pressure in thermodynamics, $[Pa]$
q_α	mass source term of phase α , $[kg/s]$
q_i^α	source term of basis species i in transport model of phase α , $[mol/s]$
Q	ion activity product
R	gas constant, $[J/K/mol]$
R_α	reaction term of phase α , $[kg/s]$
S	concentration of species in chemical module
S_j	concentration of species j in chemical module
S_α	saturation of phase α
$S_{\alpha r}$	residual saturation of phase α
T	Temperature, $^\circ C$ and K in sec.
T_i	total concentration in chemistry module
\mathbf{u}_α	Darcy's velocity of phase α
V	molar volume, $[m^3/mol]$
y_i	mole fraction in gas phase of basis species i
<hr/> Greek Symbols <hr/>	
$\alpha = \{l, g\}$	liquid/gas phase

α_{ij}	stoichiometric coefficient
β	power parameter of kinetic model
γ_i	activity coefficient
Γ	boundary domain of benchmark problem
δ	power parameter of kinetic model
λ	parameter of Brooks-Corey model
μ_α	viscosity of phase α , $[Pa \cdot s]$
ρ_α	mass density of phase α , $[kg/m^3]$
ϕ_α	source velocity of phase α , $[m/s]$
ω	porosity
Ω	global modeled domain of benchmark problem
$\ \cdot\ $	Eucliden norm

2. Geological context of the exercise

Compositional heterogeneities of hydrogen sulfide (H_2S) have been noticed in many sour gas reservoirs. H_2S occurrence is an important factor of economic depreciation. Thus, the knowledge of its content and distribution is a critical parameter when planning field development.

A possible scenario for H_2S segregation in gas reservoir stems from the preferential leaching of H_2S by underlying aquifers [23, 24]. Under conditions of pressure and temperature of typical reservoirs, H_2S and CO_2 are far more soluble than hydrocarbons, Fig. 1. A preferential leaching of H_2S (and to a lesser extent CO_2) versus CH_4 over time takes place. In the case of a deep and trapped reservoir of acid gases in contact with a bottom aquifer, Fig. 1, this differentiation is controlled by several mechanisms [23].

1. The differential solubility of gases changes the relative amounts of each gas component near the gas-water contact.
2. An active aquifer can effectively export the dissolved gases, thus enhancing dissolution on the long-term.

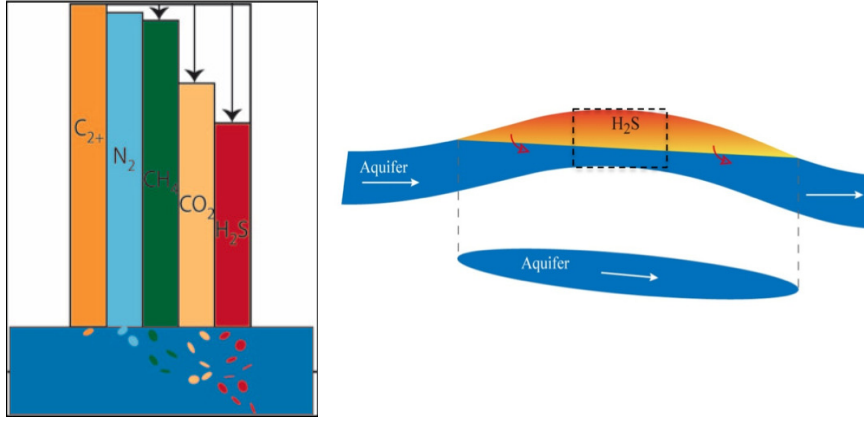


Figure 1: Schematic representation of the differential solubility of gases in water (left) and of the longitudinal section of a deep and trapped reservoir of acid gases in contact with an underlying aquifer (the dashed line stands for the localization of the simplified modeled grid).

3. Higher ratio between the length of the gas/aquifer interface and the height of the reservoir maximizes the dissolution, leading to larger amounts leached.
4. Diffusive and advective transport in the gas phase transfers the compositional anomalies farther from the gas-water contact.

This exercise is a simplified simulation of the behavior of such reservoir: the gas phase is mostly composed of methane (CH_4), with significant contents in H_2S and carbon dioxide (CO_2). In order to accurately represent the system, the simulation of several phenomena is required. A two-phase (gas, water) system description is obviously a pre-requisite. The description of a multicomponent gas phase is also compulsory to follow the evolution of the composition in the reservoir. The description of the aquifer implies the resolution of the water flow. The dissolution of the gas in the aquifer is bound to modify the local (gas) pressure, so that gaseous flow should also be represented. From a physicochemical point of view, accurate water-gas interactions can be described by several equations of state: ideal gas is of course the simplest, but high pressure high temperature conditions require the use of more complex equations. Finally, the description should include some water-rock interaction, involving the dissolved acids ($\text{CO}_2(\text{aq})$, $\text{H}_2\text{S}(\text{aq})$) and host rock minerals.

These phenomena are strongly coupled. Beyond the (two-phase) flow, transport and chemistry coupling, several interesting feedbacks should be considered. The evolution of the gas composition modifies the local gas density (H_2S is more dense than CH_4) which impacts the (compressible) gas flow. The physicochemical dissolution of gas components in the water phase removes some mass from the gas phase; this should be accounted for by resetting the pressure-volume in the flow equations, using mass-volume relationships heavily dependent on gas composition and pressure. Assuming incompressible flow and no mass exchange in the flow system for the majority of the benchmark cases allowed to simplify the system for benchmark participants and to focus on the impact of different thermo- and hydrodynamical factors [25].

Several mathematical frameworks exist to represent each of the processes described above. The most generally used governing equations and numerical methods for these problems are given in the following section.

3. Mathematical Model

3.1. Two-Phase Flow

We consider an isothermal two-phase flow with compressible gas phase and reactive multicomponent transport in porous medium. Let us introduce a two-phase flow model, which states a system of partial differential equations of mass conservation for each phase α , liquid and gas,

$$\frac{\partial \omega \rho_\alpha S_\alpha}{\partial t} + \nabla \cdot (\rho_\alpha \mathbf{u}_\alpha) = q_\alpha + R_\alpha \quad (1)$$

where the fluxes can be expressed for each phase by the Darcy-Muskat law [26, 27, 28]

$$\mathbf{u}_\alpha = -\frac{k_{r\alpha}}{\mu_\alpha} \mathbb{K} (\nabla p_\alpha - \rho_\alpha \mathbf{g}), \quad (2)$$

where $\mathbb{K} = KI$ and K is a scalar coefficient of intrinsic permeability under the isotropic conditions. The source term q_α can be also presented as

$$q_\alpha = \rho_\alpha \phi_\alpha. \quad (3)$$

To close the system we get the following constitutive relations:

$$S_l + S_g = 1, \quad (4)$$

$$p_g - p_l = p_c. \quad (5)$$

The capillary pressure p_c and relative permeability k_r are presented by empirical laws, *e.g.* Brooks-Corey model [29]:

$$\bar{S}_\alpha = \frac{S_\alpha - S_{\alpha r}}{1 - S_{lr} - S_{gr}}, \quad (6)$$

$$p_c(S_l) = p_e \bar{S}_l^{-\frac{1}{\lambda}}, \quad (7)$$

$$k_{rw}(S_l) = \bar{S}_l^{\frac{2+3\lambda}{\lambda}}, \quad (8)$$

$$k_{rn}(S_l) = (1 - \bar{S}_l)^2 (1 - \bar{S}_l^{\frac{2+\lambda}{\lambda}}). \quad (9)$$

105 The gas density is calculated using the (cubic) equations of state that will be discussed next, see section 3.3.

Numerous formalisms are proposed in the literature to solve the flow equation [30, 31, 11, 13, 32]. They can rely on substitution of several unknowns [32], so that the system is described by pressure/saturation and pressure/mole fraction in liquid phase, or the formalism of pressure/pressure [31] or pressure/mass density [11] can be applied. In the HYTEC code, the resolution is built on a water pressure/gas saturation formalism, on a Voronoi-based finite volume discretization and on a fully implicit method with analytical Jacobian. An adaptive time-stepping is controlled by the number of Newton iterations. We neglect the liquid compressibility and viscosity variation.

3.2. Multicomponent Transport

Based on the principal components formalism (see 3.4), we define a limited number of basis species N_p^{tr} and denote their total liquid mobile, immobile and gas concentration by c_i^l , \bar{c}_i and c_i^g . The liquid and gas transports are separated, the former combines liquid mobile and immobile part of species i :

$$\frac{\partial \omega S_l c_i^l}{\partial t} = \mathfrak{T}(c_i^l) + q_i^l + R_i^l(c_i^l, \bar{c}_i^l), \quad (10)$$

$$\frac{\partial \omega S_g c_i^g}{\partial t} = \mathfrak{T}(c_i^g) + q_i^g + R_i^g(c_i^g) \quad (11)$$

where operator \mathfrak{T} includes the advective flux presented by Darcy's law and the diffusive-dispersive flux [26, 33, 12]

$$\mathfrak{T}(c_i^\alpha) = \nabla \cdot (D_\alpha^{eff} \nabla c_i^\alpha - c_i^\alpha \mathbf{u}_\alpha). \quad (12)$$

Assuming the equality of longitudinal and transverse dispersion coefficients, the effective diffusion can be written:

$$D_\alpha^{eff} = \omega S_\alpha D_\alpha + a_\alpha \|\mathbf{u}\|. \quad (13)$$

The Millington-Quirk tortuosity model [34] is often used, in this case the effective diffusion takes the following form:

$$D_\alpha^{eff} = \omega^{4/3} S_\alpha^{10/3} D_\alpha + a_\alpha \|\mathbf{u}\| \quad (14)$$

In HYTEC, the transport resolution is performed by an upwind flux discretization with the harmonic weighting of diffusion. Applying a semi-implicit method for time discretization makes it possible to achieve an accurate solution: implicit Euler in time for diffusive-dispersive and Crank-Nicholson for advective flux. Other authors prefer to unify both phases and resolve implicitly fully coupled mass or mole balance for each component [35, 13, 32].

3.3. Equation Of State

An equation of state (EOS) is required to define the volumetric properties of the system: it establishes a relationship between pressure, molar volume and temperature (PVT) and allows to evaluate density, viscosity, fugacity, solubility. For any pure homogeneous fluid in the equilibrium state and for the region of single phase, this relation can be expressed by the following equation [36]:

$$f(P, V, T) = 0. \quad (15)$$

The ideal gas is the simplest form of equation of state, although it is limited to low pressure/temperature regions:

$$PV = RT \quad (16)$$

More complex EOS were developed for higher temperatures and pressures, notably the cubic equations of state which offer an accurate description of both liquid and gas phases over a wide pressure/temperature range. E.g. the Peng-Robinson EOS [37]:

$$P = \frac{RT}{V-b} - \frac{a(T)}{V(V+b)+b(V-b)} \quad (17)$$

which uses the binary interaction parameters and can handle multicomponent gases. So, applying one of the EOS provides mass density for gas mixture:

$$\rho_g = \frac{\sum_{i=1}^{N_p} y_i M_i}{V}. \quad (18)$$

130 3.4. Chemistry

Most reactive transport codes rely on the primary species formulation [15, 12, 38], where all species S_j , $j = 1, \dots, N_s$ can be expressed as a function of primary species C_i from established basis of N_p species, where $N_p < N_s$:

$$S_j = \sum_{i=1}^{N_p} \alpha_{ij} C_i \quad (19)$$

Their concentrations are constrained by the Mass-Action Law:

$$S_j = \frac{K_j}{\gamma_j} \prod_{i=1}^{N_p} (\gamma_i C_i)^{\alpha_{ij}} \quad (20)$$

using stoichiometric coefficients α_{ij} and activity coefficients γ_j . For each basis species, the total concentration T_i , $i = 1, \dots, N_p$ can be written through

$$T_i = \sum_{j=1}^{N_s} \alpha_{ji} S_j, \quad (21)$$

$$T_i - \sum_{j=1}^{N_s} \alpha_{ji} \frac{K_j}{\gamma_j} \prod_{i=1}^{N_p} (\gamma_i C_i)^{\alpha_{ij}} = 0, \quad (22)$$

the latter constitutes a system of mass conservation on C_i .

The kinetics of mineral precipitation/dissolution can be treated by a transition state theory type rate law [39]:

$$\frac{dS_j}{dt} = A_s k \left(\left(\frac{Q}{K_s} \right) - 1 \right) \prod a_i^{n_i} \quad (23)$$

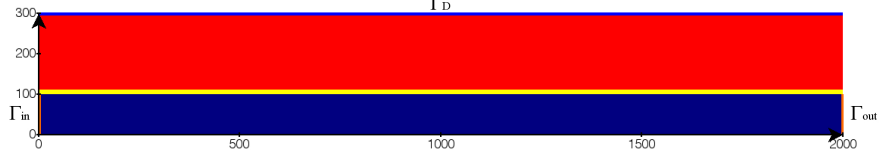


Figure 2: Geometry: Reservoir, Contact and Aquifer zones (red, yellow and blue, respectively)

where A_s is the specific surface area, k is the dissolution/precipitation rate constant and a_i is the activity of the potential catalyzing or inhibiting species. Q_s is the ion activity product defined by

$$Q_s = \frac{\gamma_j S_j}{\prod_{i=1}^{N_p} (\gamma_i C_i)^{\alpha_{ij}}}. \quad (24)$$

Mass-Action Law for all derived species at equilibrium, kinetic law for kinetically limited minerals, and mass conservation achieve a complete set of algebro-differential equations. Its resolution, at each node of the domain, yields the concentrations of all the species in the system.

4. Benchmark Description

The benchmark exercise is based on a simplified description of the geological example. The geometry is shown in Fig. 2: the 2D model is similar simplification (or a crop-out) of the general geometry Fig. 1. There are two main zones, reservoir and aquifer; the third acts as a contact/transition zone. The simulation time is 10^5 y.

Table 2 details the zone geometry and decomposition; the boundary conditions represent the average aquifer velocity 0.15 m/y at a depth 3.3 km.

$$\Gamma_{in,out} : \phi_l = (4.8e-9, 0) [m/s] \quad (25)$$

$$\Gamma_D : p_l = 1e5 + \rho_l g(3.3e3 - z) [Pa] \quad (26)$$

The geochemical and hydrodynamic description can be found in Tables 3 and 4. The initial condition *gas* is calculated for temperature 100°C and pressure 300 bar and distributed homogeneously in the Reservoir and Contact zones.

	$\{x_0, x_L; z_0, z_H\}, [m]$	S_g	p_l
Global domain Ω	$\{0, 2000; 0, 300\}$		hydrostatic
Reservoir	$\{0, 2000; 110, 300\}$	0.9	
Contact	$\{0, 2000; 100, 110\}$	0.2	
Aquifer	$\{0, 2000; 0, 100\}$	0	

Table 2: Zone decomposition

ω	0.2	
$K, [m^2]$	1e-13	
p_c	Brooks-Corey, ($p_e = 1e5, \lambda = 2, S_{lr} = 0.01$), Eq. (7)	
$k_{r\alpha}$	Brooks-Corey, ($\lambda = 2, S_{lr} = 0.01$), Eq. (9)	
$p_l, [Pa]$	$1e5 + \rho_l g(3.3e3 - z)$	
$T, ^\circ C$	100 isothermal	
α	l	g
$D_\alpha, [m^2/s]$	8e-9	5e-8
$a_\alpha, [m]$	1	–
$\mu_\alpha, [Pa \cdot s]$	1e-3	3.9e-05
$\rho_\alpha, [kg/m^3]$	997	215.5

Table 3: Hydrodynamic parameters

150 Moderate salinity, calcite, and/or siderite are added, depending on the complexity of the scenario.

Several cases were set up with increasing complexity:

1. Geochemistry: the gas composition is set for all cases, but calcite, then calcite+siderite are added for the more complex cases;
- 155 2. EOS: ideal gas or Peng-Robinson;
3. Fluid dynamics: incompressible two-phase flow is tested first using two different saturation-diffusion laws, then the full two-phase compressible flow is proposed.

Table 5 summarizes the scenarios: GCase 1 to 3, increasingly complex chem-
160 istry, ideal gas and simplified flow, FCase 1 to 7 with simple chemistry, cubic
EOS and increasing flow complexity. For sake of simplicity, the assumption of
incompressible flow excluding mass exchange in flow module is accepted in all
cases except for FCase 7, that comprehends a tight coupling with the chemistry
feedback of gas-water-rock interaction.

165 5. Numerical Results

We present some simulation results obtained using HYTEC, for scenario
FCase 1 (Table 5): advection in the whole domain, ideal gas and incompressible
flow assumptions. The focus in this paper is more on the physical description
of the simulation results: the numerical and accuracy aspects will come in a
170 second time when comparing the results from different codes.

During first $2e4$ y, the system is dominated by the gravity driven gas flow:
the gas saturation is redistributed in the reservoir, with a capillary fringe at
the bottom, and gas concentrating at the top of the formation with saturation
as high as 0.98 (Fig. 4). The system approaches to a stationary state, so the
175 capillary fringe motion decelerates afterwards. Enhanced H_2S dissolution in the
aquifer modifies the H_2S content, with a sharp decrease at the contact with
the aquifer (Fig. 3). H_2S dissolution is the strongest upwards in the aquifer,
due to fresh water renewing by the aquifer, creating a visible horizontal $H_2S(g)$
gradient (Fig. 3, 70 ky), until nearly all the H_2S is evacuated (100 ky).

180 The effect of the EOS can be observed Fig. 5: the figure illustrates results
from FCase 1 and FCase 3. Both scenarios are identical (no mineral, two-phase
flow, incompressible gas), and differ only by the EOS: ideal gas for FCase 1,
Peng-Robinson for FCase 3. The preferential leaching of H_2S is observed in
both models, but the mole fraction for FCase 3 stays significantly higher than
185 that for FCase 1. This is highlighted by the evolution of the total mass in the
system, Fig. 6. CH_4 and CO_2 loss is limited, quasi linear in time, and similar for
both EOS. On the contrary, H_2S evacuation rate is much higher (due to lower

Geochemical solution	gas			calcite		siderite/pyrrhotite
species/mineral	CO ₂ (g)	H ₂ S(g)	CH ₄ (g)	NaCl	Calcite	Siderite
y_i /concentration	0.05	0.15	0.8	0.1 molal	10 kg/l	1 kg/l

Table 4: Geochemical parameters

Part		Geochemistry			Fl.Dynamics & EOS						
Geochem.	Parameter\Case	GCCase 1/ FCCase 1	GCCase 2	GCCase 3	FCCase 2	FCCase 3	FCCase 4	FCCase 5	FCCase 6	FCCase 7	
		Gas	Calcite	Calcite Pyrrh.	$K_R = 0$ IG	$K_R \neq 0$ PR	$K_R = 0$ PR	Diff. MQ	Diff. AM	CF	
		gas	+	+	+	+	+	+	+	+	+
		calcite		+	+						
		siderite/pyrrhotite			+						
EOS	Ideal Gas	+	+	+	+						
	Peng-Robinson						+	+	+	+	
Fl.Dyn.	adv. in Reserv., $K_R \neq 0$	+	+	+		+		+	+	+	
	diffusion, Eq. (13)	+	+	+	+	+	+			+	
	diff. (MQ), Eq. (14)							+			
	adapted diff. (AM), [25]								+		
	compr. flow (CF)										+

Table 5: Benchmark Configuration

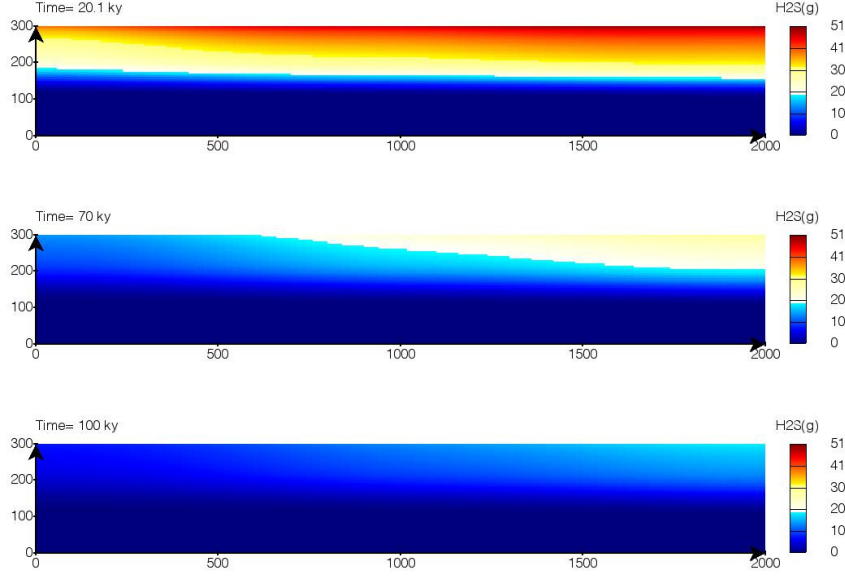


Figure 3: FCASE 1. Time evolution of $\text{H}_2\text{S}(\text{g})$ [molal]

initial content and higher solubility), and starts decreasing after 2×10^4 y, due to decreasing accessibility. More remarkable is the large difference of behavior due to the choice of EOS: ideal gas assumption largely overestimates H_2S solubility. The effect is minor on CO_2 and CH_4 , although they are not in the same direction (slight overestimation for CH_4 , slight underestimation for CO_2).

6. Conclusion

The multiphase multicomponent benchmark is proposed to the reactive transport modeling community. The exercise is based on geologically realistic foundations, and allows to explore a complex coupling involving gas-water-rock interactions. The system includes a tight connection between transport and geochemistry. The evolution of the mass/volume due to the reactive transport affects the pressure field, which leads to a strong feedback of chemistry on flow. An accurate modeling of a multicomponent gas phase is also required for this

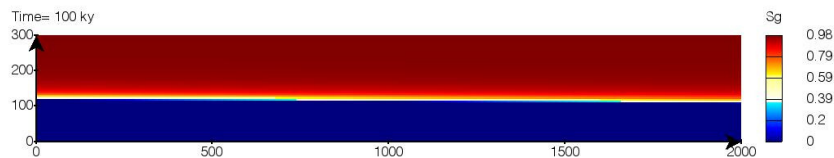


Figure 4: FCASE 1. Gas saturation. Incompressible flow.

problem. Finally, the complexity of the CO_2 and H_2S geochemistry in the host rock is of particular interest.

The benchmark problem was formulated and two sets of cases were constructed to investigate the geochemical, thermo- and hydrodynamical factors. The sensibility study showed a strong quantitative impact of the choice of equation of state, diffusion model and advection in reservoir on gas leaching.

The choice of observables for comparison between codes is not finalized yet. It will involve a combination of color maps, profiles at several rendezvous times, local evolutions at prescribed positions for gas saturation, pressure and selected species. The evolution of the total mass is also an interesting (although vastly integrative) observable. A more detailed description can be found in [25].

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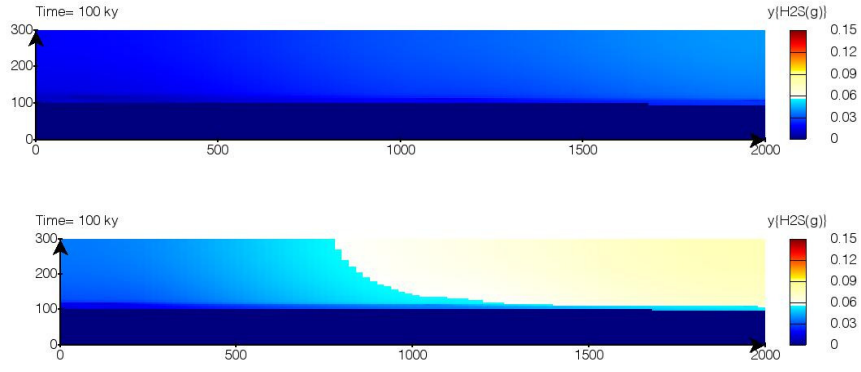


Figure 5: $\text{H}_2\text{S}(\text{g})$ mole fraction of FCASE 1 ideal gas (top) and FCASE 3 Peng-Robinson EOS (bottom)

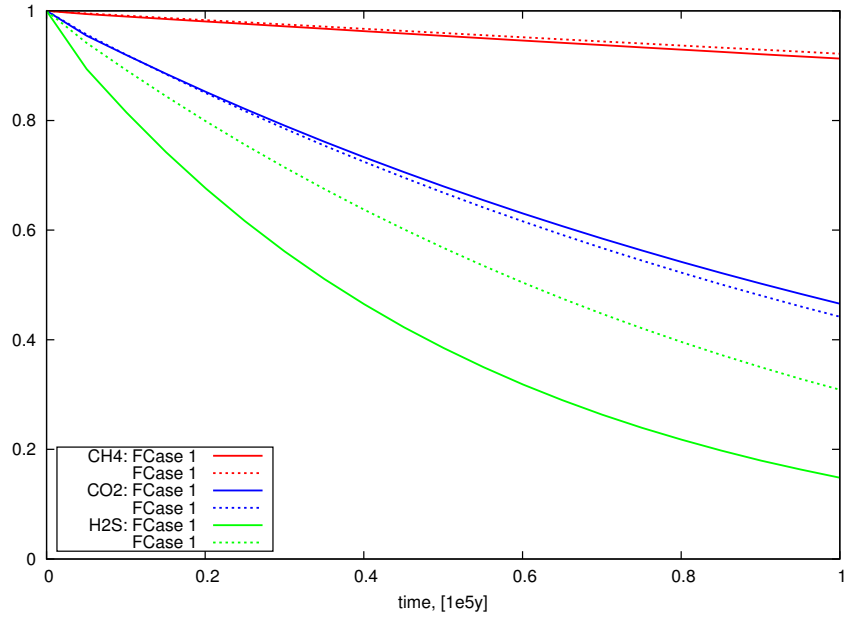


Figure 6: Time evolution of gas quantity of FCASE 1 (IG) and FCASE 1 (PR). EOS impact.

Chapter 4

Equations of State and Fluid Properties

L'écoulement et le transport multiphasiques compressibles comprennent la définition des propriétés thermodynamiques pour lesquelles les équations d'état peuvent être appliquées. Les paramètres de modèles sont calibrés en fonctions des propriétés nécessaires. Le Chapitre 4 présente une classification par type de modèle d'équation d'état. L'attention est portée sur la prédiction des propriétés volumétriques de corps purs et de systèmes binaires ; par conséquent, une analyse comparative des équations d'état est menée. Nous présentons également une liste succincte des modèles de propriétés des fluides qui sont utilisés dans les simulations numériques des Chapitres 3 et 5.

In Sec. 3.2, we have presented the coupling of miscible flow and reactive transport, where the EOS can be used for different purposes globally or/and locally. The calibration of EOS parameters and their formulations vary according to the modeled property; e.g., vapor pressure, VLE, enthalpy. Sec. 4.1 provides an overview of the equations of state classified by type of model. A comparative analysis on the prediction of volumetric properties of pure components and binary systems was carried out, Sec. 4.2. Then, Sec. 4.3 presents the selected models of fluid properties, that are used for the numerical results in Chs. 3 and 5.

4.1 Overview of equations of state

Let us give a brief survey of some existing EOS models and their applicability to different type of systems. The ideal gas law Eq. (3.1.6) is the simplest form of equation of state, although it is limited to low pressure/temperature regions. More complex EOS were developed for higher temperatures and pressures, notably the cubic equations of state which can offer an accurate description of both liquid and gas phases over a wide pressure/temperature range. Since the Peng-Robinson EOS provides a better estimation of compressibility factor, vapor pressure and liquid density than SRK, we take it as a base model for further modeling in this study.

4.1.1 Peng-Robinson model

The Peng-Robinson model (**PR78**) (Robinson and Peng, 1978) for a pure component i takes the following form:

$$P = \frac{RT}{v - b_i} - \frac{a_i(T)}{v(v + b_i) + b_i(v - b_i)}, \quad (4.1.1)$$

where the repulsive and attractive terms are expressed using the critical pressure, temperature and acentric factor, $P_{c,i}$, $T_{c,i}$, ω ,

$$a_i(T) = 0.253076587 \frac{R^2 T_{c,i}^2}{P_{c,i}} \alpha_i(T), \quad (4.1.2)$$

$$\alpha_i(T) = (1 + m_i(1 - \sqrt{T_{r,i}}))^2, \quad T_{r,i} = T/T_{c,i}, \quad (4.1.3)$$

$$m_i = \begin{cases} 0.37464 + 1.54226\omega_i - 0.26992\omega_i^2 & \text{if } \omega_i \leq 0.49, \\ 0.379642 + 1.48503\omega_i - 0.164423\omega_i^2 + 0.016666\omega_i^3 & \text{else,} \end{cases} \quad (4.1.4)$$

$$b_i = 0.0777960739 \frac{RT_{c,i}}{P_{c,i}}. \quad (4.1.5)$$

The difference between PR76 and PR78 is m_i parameter, namely the ω condition. In the mixture, the parameters a and b of PR78 are calculated through the combining rule for the cross parameter a_{ij} and the quadratic mixing rules for a and b :

$$a = \sum_i \sum_j x_i x_j a_{ij}, \quad (4.1.6)$$

$$a_{ij} = \sqrt{a_i a_j} (1 - k_{ij}), \quad (4.1.7)$$

$$b = \sum_i x_i b_i, \quad (4.1.8)$$

where k_{ij} are the binary interaction parameters. By using Eqs. (4.1.6)-(4.1.8), the cubic EOS yields the phase behavior of mixtures of nonpolar and slightly polar compounds and it gives a less adequate VLE description for the strongly polar systems, such as alcohols and water. A large spectrum of methods and modifications now exists to improve EOS models. We first discuss the impact of modified α and b functions on the VLE prediction. Then the Volume Translation (VT) method and the derivation of mixing rules from the excess Gibbs energy will be concisely described below. The generalization of two/three constant EOS and also the derivation of main EOS except can be found in Schmidt and Wenzel (1980), Ji and Lempe (1998), Michelsen and Mollerup (2007).

4.1.2 α function: vapor pressure

In order to give a better description of vapor pressure and VLE, the modified α function were proposed by representing it in the polynomial or/and exponential form, by applying the different formulation for sub- and supercritical regions, $T_r \leq 1$ and $T_r > 1$. Considering that the attractive forces are expressed through the a parameter and consequently, through the α function, the latter should obey the next requirements: $0 < \alpha(T) < C$, C is the positive real constant, $\forall T$; $\alpha(T_c) = 1$. Moreover, the interaction between molecules weakens with the increase of temperature, so $\lim_{T \rightarrow \infty} \alpha(T) = 0$. Note that the PR78 α function Eq. (4.1.2) does not fulfill this condition. Finally, to keep the consistency of the whole thermodynamic system, $\alpha(T) \in C^2(T)$.

Twu et al. (**TPR**) generalized the α function, switched in the sub- and supercritical regions:

$$\alpha(T_r) = \alpha^{(0)}(T_r) + \omega \left(\alpha^{(1)}(T_r) - \alpha^{(0)}(T_r) \right), \quad (4.1.9)$$

$$\alpha^i(T_r) = T_r^{N_i(M_i-1)} \exp L_i (1 - T_r^{N_i M_i}), \quad i = 0, 1, \quad (4.1.10)$$

where L , M , N are the constants that can be either defined for each compound (Twu et al., 1995) or generalized for hydrocarbon fluids by using two points $P_r = P/P_c = 1$ and $T_r = 0.7$. Besides, the generalized parameters differ according to the critical temperature point, see Table 4.1. The vapor pressure for light and heavy hydrocarbons was predicted with 3.3% ADD compared to 12.1% of PR76 (using Eq. (4.1.4)) from the triple point to the critical one. The L , M , N parameters were later adjusted to a wider range of components

Table 4.1 – Parameters for the TPR EOS model: L , M , N parameters of Eq. (4.1.10) for hydrocarbon fluids (Twu et al., 1995); (L), (M), (N) parameters, generalized for alkanes, aromatics, ketons, alcohols, refrigerants and gases at $T_r \leq 1$ (Ahlers and Gmehling, 2002b).

	$T_r \leq 1$		$T_r > 1$	
	$\alpha^{(0)}$	$\alpha^{(1)}$	$\alpha^{(0)}$	$\alpha^{(1)}$
L	0.125283 (1.511442)	0.511614 (0.567879)	0.401219	0.024955
M	0.911807 (2.788270)	0.784054 (0.774270)	4.963070	1.248089
N	1.948150 (0.161590)	2.812520 (2.575067)	-0.2	-8.0

at $T_r < 1$, see Table 4.1. In the supercritical region, it was fitted in regard to the solubility of hydrogen and methane in hydrocarbon liquids.

Gasem et al. (**GPR**) proposed a modified α function to improve the vapor pressure prediction of pure components without switching it for sub- and near critical conditions as was given in Mathias and Copeman (1983):

$$\alpha(T) = \exp \left((A + BT_r) \left(1 - T_r^{C+D\omega+E\omega^2} \right) \right), \quad (4.1.11)$$

where $A - E$ are the generalized constants. The GPR model displayed the vapor pressure with 1.1% AAD for 28 pure compounds (3.7 AAD for water) and the heat capacity with 3% AAD for methane, propane and nitrogen for sub- and supercritical zones. AAD is defined as $100/N \sum_{i=1}^N |X_{calc} - X_{exp}|/X_{exp}$.

Coquelet et al. (**CPR**) combines both the exponential and polynomial form of α function

$$\alpha(T) = \begin{cases} \exp [c_1 (1 - T_r)] \left[1 + c_2 (1 - \sqrt{T_r})^2 + c_3 (1 - \sqrt{T_r})^3 \right]^2 & \text{if } T_r \leq 1, \\ \exp [c_1 (1 - T_r)] & \text{else,} \end{cases} \quad (4.1.12)$$

where c_i , $i = 1, 2, 3$ are the fitted parameters that have the polynomial form of ω : $\sum_{k=0}^2 c_{i,k} \omega^k$, $c_{i,k}$ are the generalized constants, calculated over the selected compounds through the objective function of pressure. Using $c_{i,k}$ estimated for the system of 22 pure compounds, the vapor pressure was predicted with 1.2% AAD (5% AAD for water) in contrast to 2.1% of PR76, while the fitting over 8 pure compounds provided 0.8% AAD (0.4% AAD for water).

Table 4.2 – Comparison of PR78, SRK, GPR, HPR. AAD% of the vapor pressure and liquid density of Ar, CO₂, N₂, SO₂, O₂, H₂S, H₂O and overall 49 pure compounds, extracted from Haghtalab et al. (2011).

	vapor pressure				liquid density			
	PR78	SRK	GPR	HPR	PR78	SRK	GPR	HPR
Ar	0.31	1.64	1.08	1.05	10.34	3.62	10.44	3.63
CO ₂	0.71	0.39	1.84	0.90	3.16	10.88	3.12	2.42
N ₂	0.67	1.19	0.21	0.42	9.79	3.46	9.84	3.45
SO ₂	2.39	2.08	2.87	1.94	2.33	12.98	2.37	2.72
O ₂	1.57	1.54	0.41	0.45	10.61	3.37	10.69	3.60
H ₂ S	2.12	1.17	1.83	1.19	7.24	5.22	7.23	2.26
H ₂ O	3.33	5.91	3.94	4.20	18.44	27.79	18.48	17.50
Overall	1.6	1.7	1.5	1.4	7.2	11.7	7.2	4.8

4.1.3 α and b functions: vapor pressure and liquid molar volume

Haghtalab et al. (**HPR**) changed both the α and the co-volume b functions to improve the estimation of the vapor pressure and the volumetric properties:

$$\alpha(T) = \exp\left(1 - c_1^{\ln T_r}\right), \quad (4.1.13)$$

$$b = 0.0778 \frac{RT_c}{P_c} \beta(T), \quad (4.1.14)$$

$$\beta(T) = 1 + c_2(1 - T_r), \quad (4.1.15)$$

where c_1 and c_2 are the fitted parameters in the polynomial form of ω : $\sum_{k=0}^2 c_{i,k} \omega^k$, $c_{i,k}$ are the given constants, that were calculated using the saturated vapour pressure experimental data of 53 pure compounds and the liquid density data of 49 pure compounds.

The PR78, SRK, GPR, HPR models were compared on vapor pressure, liquid density and vapor volume of pure compounds. In Table 4.2, we present the results of the selected pure components from Haghtalab et al. (2011), where the overall AAD is given for 49 pure components. As expected, PR78 provides slightly better estimation of vapor pressure than SRK, then the refined results are obtained by GPR and HPR, since both of them modified

the α function and fitted it to the vapor pressure calculation. The additional adjustment on the co-volume parameter was made in HPR in contrast to GPR and HPR, that yields to the improved liquid density. However, the water density AAD is still about 18%.

4.1.4 Volume translation

The volume translated (**VT**) method is another way to impact on the volumetric properties. P  neloux et al. introduced the *pseudo volume*

$$\tilde{V} = V + \sum_i c_i n_i, \quad (4.1.16)$$

where V is the volume, n_i is the number of moles of component i , c_i is the constant translation parameter or the correction of molar volume, which can be applied as an adjusted parameter for the volumetric properties. If the VLE is solved on *pseudo volume*, the *pseudo partial molar volume* should be introduced

$$\tilde{v}_i = \left(\frac{\partial \tilde{V}}{\partial n_i} \right)_{T,P,n_j,j \neq i} = v_i + c_i, \quad (4.1.17)$$

and consequently, the *pseudo fugacity coefficients* is derived from the fugacity coefficient definition

$$\ln \tilde{\phi}_i = \int_0^P \left(\frac{\tilde{v}_i}{RT} - \frac{1}{P} \right) dP = \ln \phi_i + \frac{c_i P}{RT}. \quad (4.1.18)$$

If one writes the $\phi - \phi$ phase equilibria by using the pseudo fugacity coefficients, the second term of the RHS of Eq. (4.1.18) appears symmetrically, that confirms the preserved consistency of the method. In the $\phi - \gamma$ system, it can be directly integrated in the Poynting factor Eq. (3.1.10) inasmuch as the impact of the translation parameter is expressed by the separate term.

P  neloux et al. applied the VT method to SRK with the three formulations for c , among which the volume correction is the function of the critical compressibility factor z_c : $c(z_c) = 0.40768(RT_c/P_c)(0.29441 - z_c)$. All three models were tested on 233 compounds and exhibited the conspicuous results. Even though the $c(z_c)$ approach was the second best due to the lack of the precise values of z_c , it showed overall 3% AAD against 11% of SRK and 5% of PR76.

Thereafter, Ahlers and Gmehling (2001) proposed the volume translated PR model (VTPR) and temperature dependent VTPR model (T-VTPR). Basing on PR78, **VTPR**

model is formulated by following

$$P = \frac{RT}{\tilde{v} - b_i} - \frac{a_i(T)}{\tilde{v}(\tilde{v} + b_i) + b_i(\tilde{v} - b_i)}, \quad (4.1.19)$$

where $\tilde{v} = v + c$ is the pseudo molar volume defined in Eq. (4.1.17). Similar to P  neloux et al. (1982), the correction volume function c is defined through the critical compressibility factor:

$$c = \delta \frac{RT_c}{P_c} (\chi - z_c), \quad (4.1.20)$$

where δ and χ are the constant parameters fitted for each compound at the temperature range $T_r < 0.8$, the data for 44 pure compounds can be found in Ahlers and Gmehling (2001). After optimization procedure, the average δ and the linear correlation between χ and z_c were defined as $\delta = 0.252$ and $\chi = 2.5448z_c - 0.4024$, that leads to the generalized formulation of c (**gVTPR**):

$$c(z_c) = 0.252 \frac{RT_c}{P_c} (1.5448z_c - 0.4042). \quad (4.1.21)$$

For $T_r < 0.8$, the molar volume v is overestimated by VTPR, it is therefore adjusted by $c(T)$ function which is derived from its value at the critical point and the temperature adjusting term (**T-VTPR**):

$$v = v_{VTPR} - c(T) = v_{VTPR} - c_c \zeta(T_r), \quad (4.1.22)$$

$$c_c = (0.3074 - z_c) \frac{RT_c}{P_c}, \quad (4.1.23)$$

$$\zeta(T_r) = \frac{0.35}{0.35 + (\eta |T_r - \alpha(T_r)|)^\gamma}, \quad (4.1.24)$$

where η and γ are the constant parameters fitted for each compound. Then they were also generalized as the functions of z_c on the same compounds except water, chloroform and methanol, that do not lie on the η curve as other compounds (**gT-VTPR**):

$$\eta = -74.458z_c + 26.966, \quad (4.1.25)$$

$$\gamma = 246.78z_c^2 - 107.21z_c + 12.67. \quad (4.1.26)$$

In order to establish the generalized models gVTPR and gT-VTPR, the critical compressibility factor z_c is thus required in addition to the standard P_c , T_c and ω .

The comparison of PR78, SRK, VTPR, gVTPR, T-VTPR, gT-VTPR models on the liquid density in the sub- and near critical regions is presented in Table 4.3, the results

Table 4.3 – Comparison of PR78, SRK, VTPR, gVTPR, T-VTPR, gT-VTPR. AAD% of the liquid density of Ar, CO₂, N₂, CH₄, H₂O and overall 44 pure compounds, extracted from Ahlers and Gmehling (2001).

	PR78	SRK	VTPR	gVTPR	T-VTPR	gT-VTPR
Ar	8.87	5.92	4.12	4.21	2.03	2.08
CO ₂	6.03	14.26	6.36	6.85	0.87	0.88
N ₂	8.50	7.08	4.70	4.73	2.33	2.38
CH ₄	10.09	5.14	3.08	3.63	2.30	2.37
H ₂ O	21.22	29.98	10.81	12.14	1.20	3.64
Overall	6.7	13.1	3.0	4.1	2.0	2.5

are taken from Ahlers and Gmehling (2001). The dominant position of (g)T-VTPR is distinct and these models can be advantageously applied for the liquid density estimation. Nevertheless, the further usage of these models at high pressure and at the critical point leads to unphysical values, because isotherms intersect due to the introduced additional volume shift. Hence, the reliability strictly depends on the PT region and on the purpose of their usage. The (g)VTPR models provide less accurate results than (g)T-VTPR, but still better in total than PR78 and SRK.

4.1.5 Excess Gibbs energy and mixing rules

The mixing rules for a and b defined by Eqs. (4.1.6) and (4.1.8) can be differently derived. Assuming that the excess volume is zero $V^E = 0$ and consequently, the excess Gibbs energy equals the excess Helmholtz energy $G^E = A^E$, the latter can be derived from the equation of state and from the mixing rules, that was firstly made for van der Waals EOS. Conversely, applying the excess Gibbs energy model and introducing it in the EOS, the mixing rules can be found, called the G^E mixing rules. It improves the VLE estimation of mixtures in sub-/critical regions and makes the parameters a and b temperature dependent. The G^E models, assuming G^E at infinite pressure or at zero pressure, imply the use of UNIFAC method (Fredenslund et al., 1975) for G^E and the activity coefficients calculations. This approach was applied for VTPR model (Ahlers and Gmehling, 2002a,b) in combination

with the GCM and the generalized α function of TPR (Twu et al., 1995).

Another method was discovered by Peneloux et al. (1989) through the van Laar equation and the standard mixing rules, that yields

$$\frac{a}{b} = \sum_i x_i \frac{a_i}{b_i} - \frac{A^E}{C_{EOS}}, \quad (4.1.27)$$

$$A^E = \frac{1}{2} \frac{\sum_i \sum_j b_i b_j x_i x_j E_{ij}}{\sum_i x_i b_i}, \quad (4.1.28)$$

$$k_{ij} = \frac{E_{ij} - (\delta_i - \delta_j)^2}{2\delta_i \delta_j}, \quad (4.1.29)$$

$$\delta_i = \frac{\sqrt{a_i(T)}}{b_i}, \quad (4.1.30)$$

where to be rigorous, A^E is the excess Helmholtz, E_{ij} are the energy binary parameters. This method was employed to PR78 in Jaubert and Mutelet (2004), called the predictive PR78 (**PPR78**). This evolves into the temperature dependent binary energy interaction parameters $E_{ij}(T)$ and hence, $k_{ij}(T)$ that are expressed by the GCM. Considering that each molecule i is composed of $N_{g,i}$ groups (e.g., CH₄, CH₃, CH₂) and that the group k is presented twice, then one can define α_{ik} as a fraction of group k , $\alpha_{ik} = 2/N_{g,i}$. Knowing the distribution $\{\alpha_{ik}\}$, $\{\alpha_{jk}\}$ of two molecules i and j , the binary interaction parameters k_{ij} are given by

$$E_{ij}(T) = -\frac{1}{2} \sum_{k=1}^{N_g} \sum_{l=1}^{N_g} (\alpha_{ik} - \alpha_{jk})(\alpha_{il} - \alpha_{jl}) A_{kl} \left(\frac{298.15}{T} \right)^{\frac{B_{kl}}{A_{kl}} - 1}, \quad (4.1.31)$$

where N_g is the total number of groups, A_{kl} and B_{kl} are the group constants, defined in Qian et al. (2013) for 20 groups, covering the petroleum fluids and natural gases. $\{A_{kl}\}$ and $\{B_{kl}\}$ form the symmetric matrices with the zero diagonals. $a_i(T)$ and b_i correspond to those of PR78, Eqs. (4.1.2) and (4.1.5), respectively. The method allowed to improve the prediction of the phase equilibria, the dew, bubble and even critical curves for binary mixtures and synthetic fluids.

4.1.6 Cubic plus association

In order to take into account the hydrogen bonding and not only spherical molecules, the cubic EOS are combined with the Wertheim's term (Kontogeorgis et al., 1996; Shinta and Firoozabadi, 1995; Hajiwi et al., 2015) that allows to describe the thermodynamic properties

of pure associating components (e.g., water, alcohols) and its mixtures. Unifying the cubic PR78 and associative term results in the CPA-PR (Hajiw et al., 2015):

$$P = \frac{RT}{v - b_i} - \frac{a_i(T)}{v(v + b_i) + b_i(v - b_i)} - \frac{1}{2} \frac{RT}{v} \left(1 + \rho \frac{\partial \ln g}{\partial \rho} \right) \sum_{i=1}^N x_i \sum_{A_i} (1 - X_{A_i}), \quad (4.1.32)$$

where g is the modified radial distribution function of (Kontogeorgis et al., 1996), ρ is the molar density, x_i is the mole fraction of the component i , X_{A_i} is the mole fraction of the molecule i not bonded at a site A. X_{A_i} depends on two parameters additional to PR78 – the association energy $\varepsilon^{A_i B_i}$ and the association volume $\beta^{A_i B_i}$, that can be neglected for the non-associative components. The model demonstrated promising results of the volumetric properties on the light, medium hydrocarbons-water systems and also CO₂ and H₂S water mixtures¹ but at cost of computational time making it not suitable for the reservoir modeling.

The choice of the EOS model should be made carefully when one looks both for the phase equilibria of pure compounds and complex mixtures at sub-/critical regions and for the volumetric properties of both phases. The conventional PR78 can be applied for the solubility and gas density evaluation. The models with modified α function (TPR, CPR, GRP, HPR) provide rather better solubility results than those of PR78. The PR78 model with the binary interaction parameters predicted by PPR78 is able to reproduce the phase equilibria of mixtures, its dew, bubble and critical points. The improvement of liquid density can be expected by modifying the mixing rule of the covolume parameter. Although, the volume translation methods ((g)VTPR, (g)T-VTPR) significantly increase the liquid density accuracy both of pure compounds and of mixtures, Sec. 4.2, the usage at high pressures should be avoided (Ahlers and Gmehling, 2001). The CPA-PR predicts the water systems, but it is not acceptable near the critical point. In order to improve the vapor pressure estimation, the α function can be modified. In MMF modeling, applying the different EOS models for each sub-problem is possible but entails inconsistencies. Therefore, depending on the principal goal of the modeling and PT region, the corresponding equations of state should be chosen.

¹C. Coquelet, M. Hajiw, personal communication, 2015: for instance, the AAD of water density is 2.7% at [273.15, 643.15] K and < 1% at [273.15, 573.15] K.

The next modified PR78 models were implemented in HYTEC: the TPR model (the modified α function) in combination with the (g)VTPR and (g)T-VTPR models (the liquid density correction) and the temperature dependent k_{ij} parameters defined by the PPR78 model.

4.2 EOS comparative study on volumetric properties

4.2.1 Pure compounds

The EOS volumetric properties study of pure compounds was fulfilled for

- CO_2 , H_2S , CH_4 at 100°C , 100 – 350 bar ($T_r > 1$);
- CO_2 , Ar, O_2 , N_2 , SO_2 at 50°C , 100 – 250 bar ($T_r > 1$ except SO_2 for which $T_r = 0.8$).

The PR78, TPR, VTPR, T-VTPR, GERG-2008 models and NIST database were chosen at this stage of the study. The GERG-2008 EOS (Kunz et al., 2007) was employed by the thermodynamic properties software REFPROP (Lemmon et al., 2013). The GERG-2008 results correspond to the NIST data for the pure compounds, so they are omitted.

CO_2 at 50 and 100°C , 20 – 350 bar

Fig. 4.1 presents the density of gas and supercritical CO_2 at 50 and 100°C and to 20 from 350 bar, provided by the PR78, TPR, VTPR, T-VTPR and NIST database. In general, the density is underestimated except the T-VTPR at $P > 160$ bar and the TPR at $P > 260$ bar.

Table 4.4 shows the AAD of the relevant EOS models: for the pressure range 20 – 300 bar and $T = 50^\circ\text{C}$, the PR78 and TPR yield about 3%, the VT models 4%; the accuracy of all 4 models improves at 100°C . The AAD varies significantly with the pressure range of interest. So, at 50°C , the **PR78**, **TPR** and **T-VTPR** models can be applied for the pressure range 100 – 250 bar, yielding 3 – 4% AAD. However, we should note that the AAD of PR78 and TPR exceed 8% at 120 – 130 bar. At 100°C , **all four models** give acceptable results within 4% AAD.

H_2S at 100°C , 100 – 350 bar

Fig. 4.2 shows the supercritical density of H_2S at 100°C and 100 – 350 bar modeled by PR78, TPR, VTPR, T-VTPR and GERG-2008 (NIST). There is an obvious discrepancy between the EOS results: the T-VTPR curve is close to that of NIST near $P_r = 1$, then the deviation grows with the increasing pressure and reaches 19% ADD. The PR78 and TPR with 2% AAD are preferable for the range 100 – 150 bar, while the VTPR provides 1% for higher pressures 250 – 350 bar, see Table 4.2.

CH_4 at 100°C , 100 – 350 bar

All models give a relevant density of supercritical CH_4 at the chosen TP conditions,

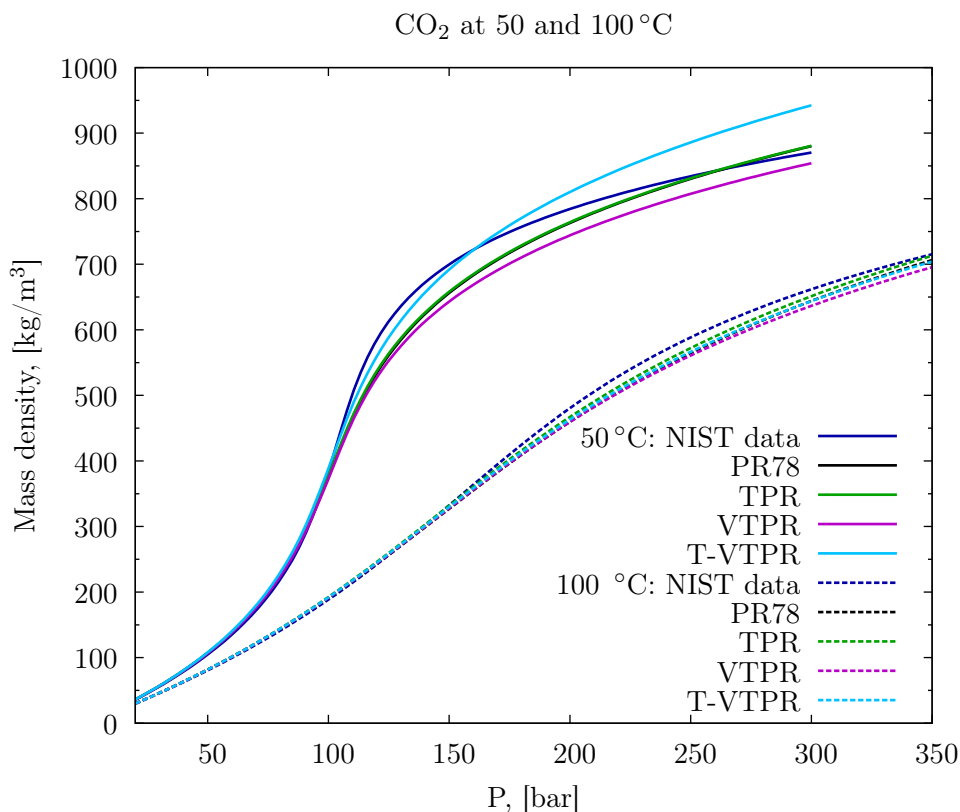


Figure 4.1 – Density of pure CO_2 at 50 and 100 °C and 20 – 350 bar provided by PR78, TPR, VTPR, T-VTPR and NIST data.

Fig. 4.3, within 3% of AAD Table 4.6. The volume translation models maintain about 1% AAD at the whole pressure range.

CO_2 , H_2S , CH_4 at 100 °C, 100 – 350 bar: At high temperature and pressure conditions, 100 °C and 250–350 bar, the **VTPR** and **PR78** models provide the acceptable gas/supercritical density (4.6% AAD at maximum) of the pure compounds CO_2 , H_2S , CH_4 . In the T-VTPR model (Ahlers and Gmehling, 2001), H_2S was not taken into account. Thus we used the generalized formulation for the temperature dependent VT parameter $c(T)$ Eq. (4.1.22), resulting in up to 19% AAD. The H_2S density can be well described (0.4% AAD) by adjusting $\eta = 5$ and $\gamma = -0.2$, Eqs. (4.1.25) and (4.1.26) respectively. Nevertheless, considering the generalized form of $\gamma > 0 \forall z_c$ Eq. (4.1.26), H_2S is out of the model conception. At 100 °C and 100 – 150 bar, the VTPR model reaches 7% AAD, hence the **PR78** is preferable. On the whole pressure range 100 – 350 bar, the **PR78** is recommended for the volumetric properties evaluation of the selected pure compounds.

N₂, O₂, Ar, SO₂ at 50 °C, 100 – 250 bar: Figs. 4.4 – 4.7 demonstrate the predicted densities by the PR78, TPR, VTPR, T-VTPR and GERG-2008 models. The TPR model yields the maximum 3.4% AAD (N₂), the best evaluation is performed by the VTPR, 0.9% for all 4 compounds, see Tables 4.7 – 4.10. The PR78 model overall provides 2%. The **PR78**, **VTPR** and **T-VTPR** allow to model the density of pure compounds with the deviation up to 2%.

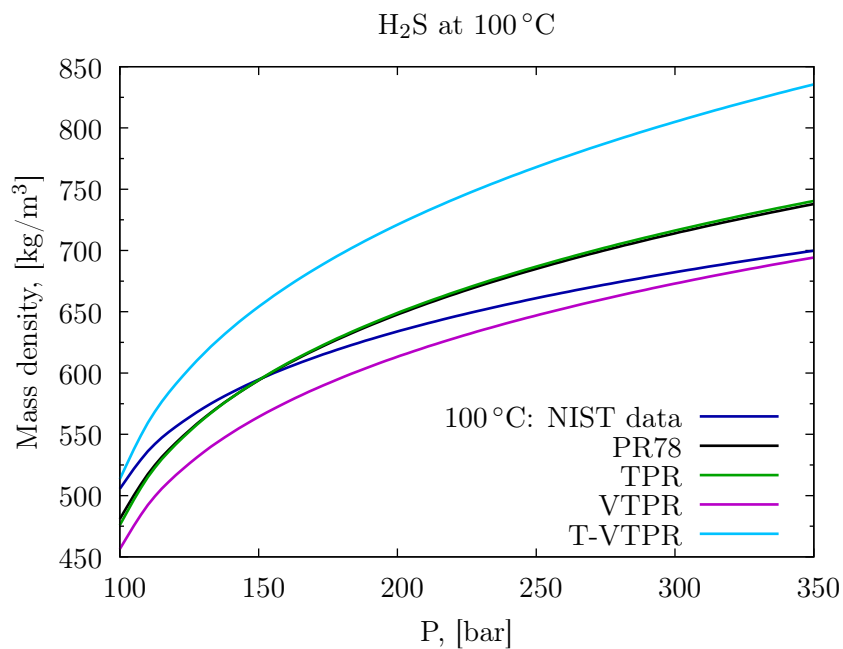


Figure 4.2 – Density of pure H_2S at 100°C and 100 – 350 bar provided by PR78, TPR, VTPR, T-VTPR and NIST data.

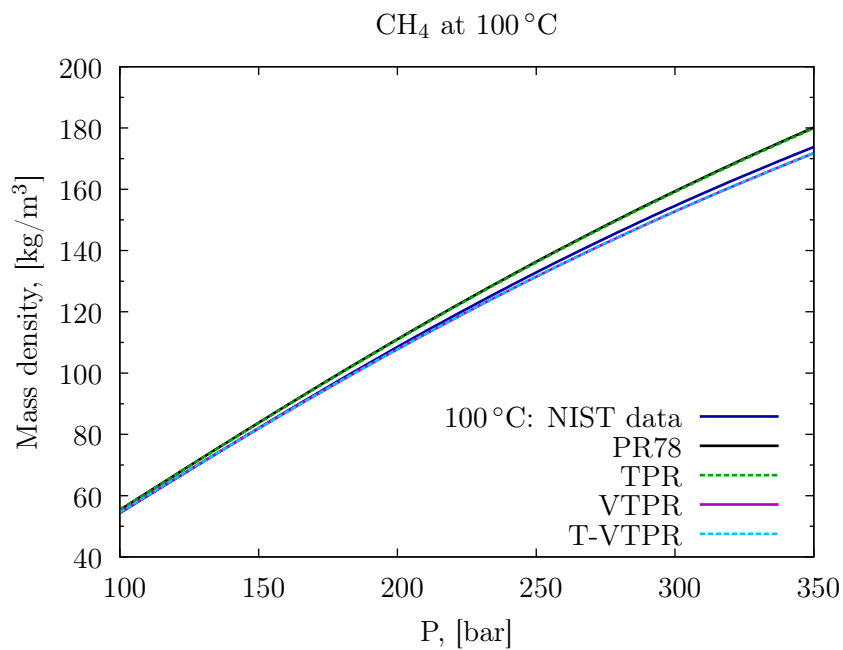


Figure 4.3 – Density of pure CH_4 at 100°C and 100 – 350 bar provided by PR78, TPR, VTPR, T-VTPR and NIST data.

Table 4.4 – Comparison of PR78, TPR, VTPR, T-VTPR. AAD% of the gas and supercritical density of CO₂ at 50 and 100 °C, 20 – 350 bar.

$T = 50\text{ }^{\circ}\text{C}$				
P, bar \	PR78	TPR	VTPR	T-VTPR
20 – 300	2.9	2.8	4.3	3.9
100 – 120	6.2	5.5	7.0	3.0
100 – 250	4.1	3.9	6.1	3.1

$T = 100\text{ }^{\circ}\text{C}$				
P, bar \	PR78	TPR	VTPR	T-VTPR
100 – 350	2.7	1.8	3.2	2.5
100 – 150	0.5	0.7	0.5	0.5
250 – 350	2.7	1.6	3.8	2.7

Table 4.5 – Comparison of PR78, TPR, VTPR, T-VTPR. AAD% of the supercritical density of H₂S at 100 °C, 100 – 350 bar.

P, bar \	PR78	TPR	VTPR	T-VTPR
100 – 350	3.2	3.4	3.4	13.8
100 – 150	2.2	2.5	7.0	6.5
250 – 350	4.6	4.9	1.4	17.9

Table 4.6 – Comparison of PR78, TPR, VTPR, T-VTPR. AAD% of the supercritical density of CH₄ at 100 °C, 100 – 350 bar.

P, bar \	PR78	TPR	VTPR	T-VTPR
100 – 350	2.6	2.5	0.8	0.8
100 – 150	1.1	1.1	0.1	0.1
250 – 350	3.1	2.9	1.2	1.2

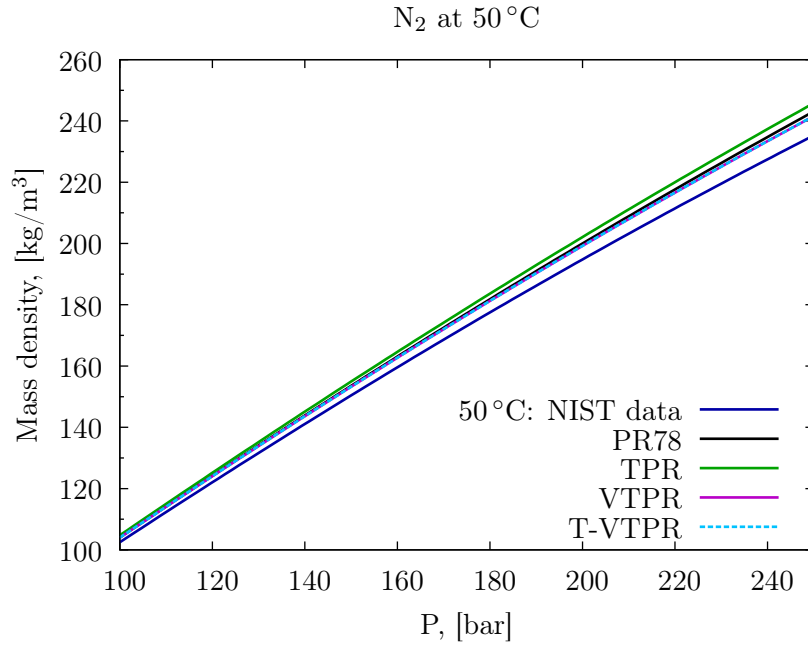


Figure 4.4 – Density of pure N_2 at $50^\circ C$ and 100 – 250 bar provided by PR78, TPR, VTPR, T-VTPR and NIST data.

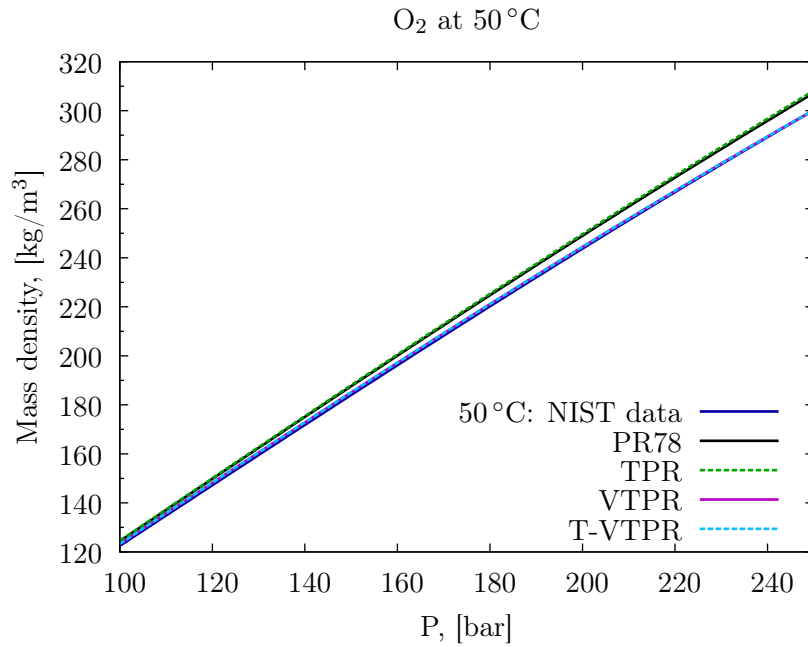


Figure 4.5 – Density of pure O_2 at $50^\circ C$ and 100 – 250 bar provided by PR78, TPR, VTPR, T-VTPR and NIST data.

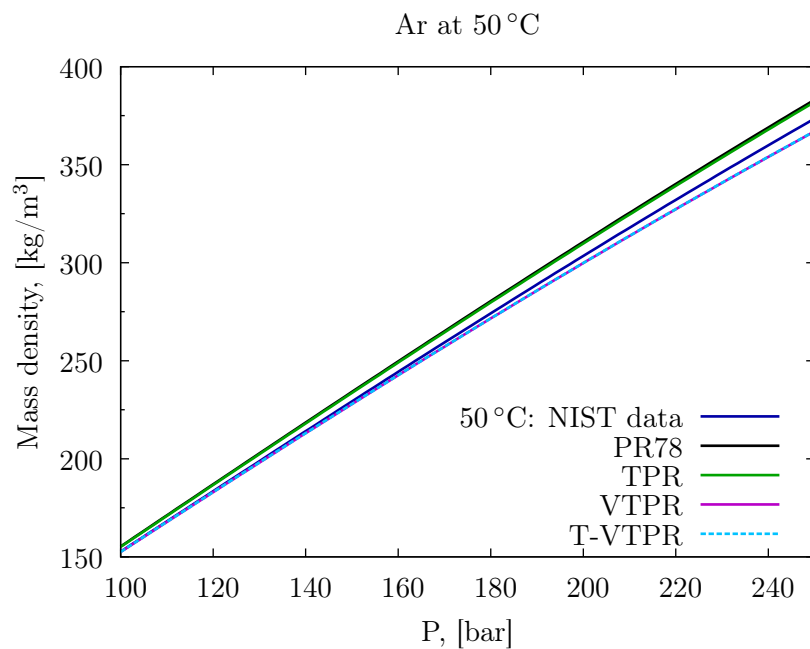


Figure 4.6 – Density of pure Ar at 50 °C and 100 – 250 bar provided by PR78, TPR, VTPR, T-VTPR and NIST data.

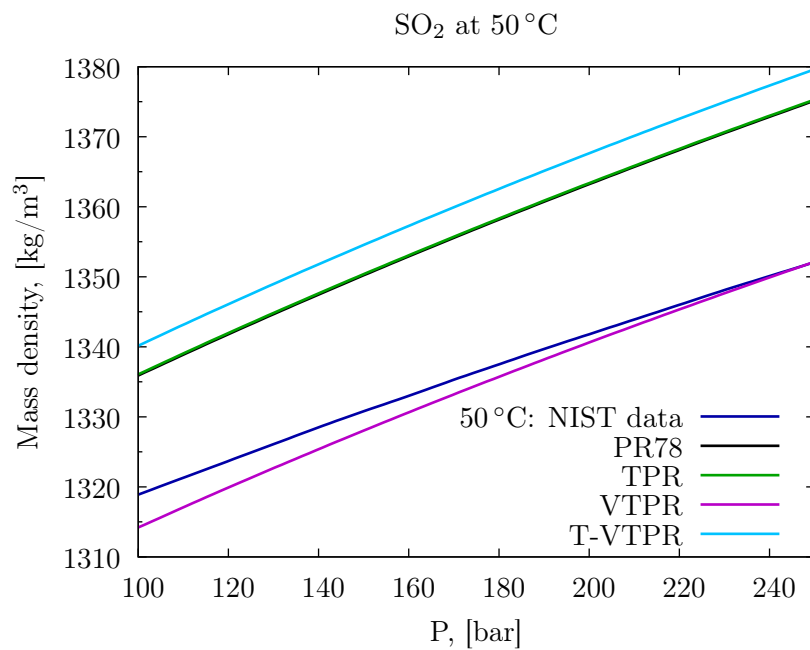


Figure 4.7 – Density of pure SO₂ at 50 °C and 100 – 250 bar provided by PR78, TPR, VTPR, T-VTPR and NIST data.

Table 4.7 – Comparison of PR78, TPR, VTPR, T-VTPR. AAD% of the supercritical N₂ density at 50 °C, 100 – 250 bar.

P, bar \	PR78	TPR	VTPR	T-VTPR
100 – 120	1.6	2.4	1.6	1.6
100 – 250	2.4	3.4	2.1	2.1

Table 4.8 – Comparison of PR78, TPR, VTPR, T-VTPR. AAD% of the supercritical O₂ density at 50 °C, 100 – 250 bar.

P, bar \	PR78	TPR	VTPR	T-VTPR
100 – 120	1.6	1.8	0.6	0.6
100 – 250	2.0	2.2	0.4	0.4

Table 4.9 – Comparison of PR78, TPR, VTPR, T-VTPR. AAD% of the supercritical Ar density at 50 °C, 100 – 250 bar.

P, bar \	PR78	TPR	VTPR	T-VTPR
100 – 120	1.8	1.6	0.2	0.2
100 – 250	2.2	2.0	0.9	0.9

Table 4.10 – Comparison of PR78, TPR, VTPR, T-VTPR. AAD% of the liquid SO₂ density at 50 °C, 100 – 250 bar.

P, bar \	PR78	TPR	VTPR	T-VTPR
100 – 120	1.3	1.3	0.3	1.7
100 – 250	1.5	1.5	0.2	1.8

4.2.2 Binary system $\{\text{CO}_2 + \text{H}_2\text{O}\}$: (T-)VTPR vs. experimental data

We present the AAD of the binary system liquid density predicted by VTPR models, because only pure compounds were considered in Ahlers and Gmehling (2001). The linear mixing rule for the volume translation c was employed. The reference 29 experimental data of $\{\text{CO}_2 + \text{H}_2\text{O}\}$ presented in Li et al. (2004) covers the pressure range 30 – 290 bar at 59 °C, hence the CO_2 reduced temperature $T_r = 1.1$. The results are given in Fig. 4.8. The VTPR model underestimates the liquid density providing 3.6% AAD while the T-VTPR overestimation is of 2.5% AAD. The accuracy of VTPR models for the binary system is of the same magnitude as that for pure compounds Table 4.3.

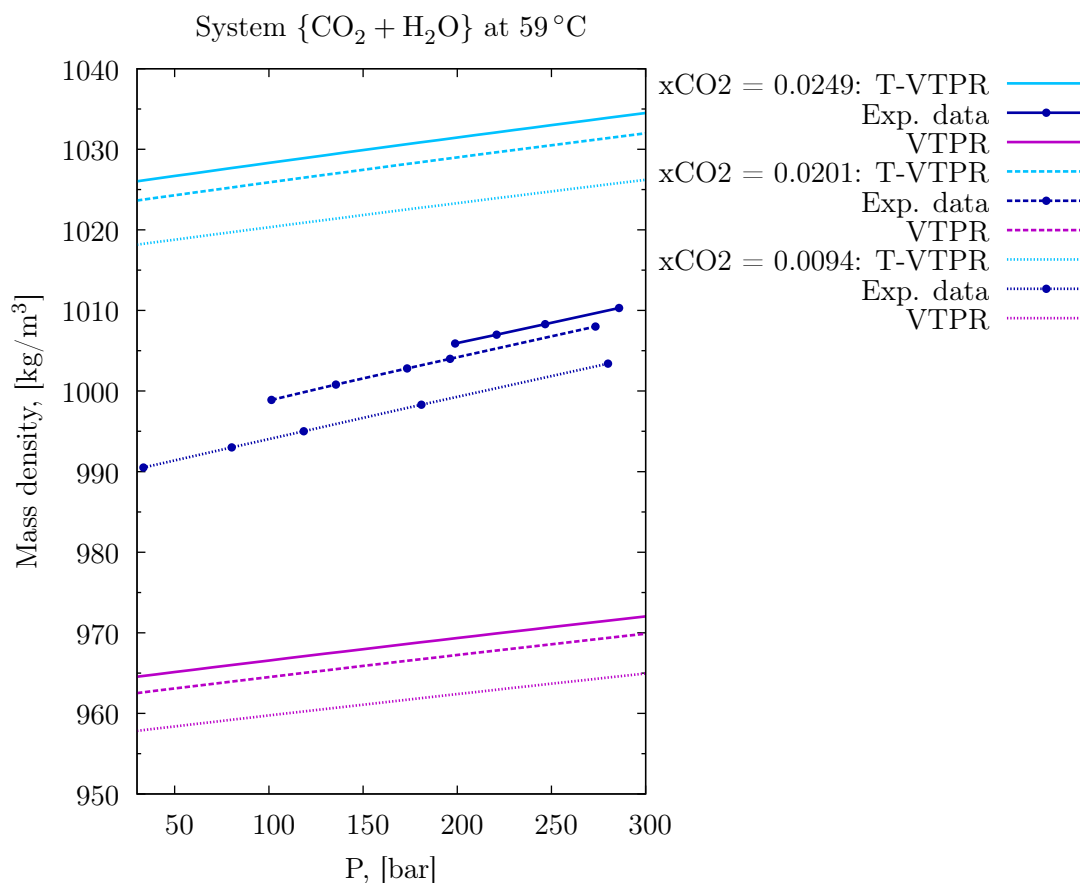


Figure 4.8 – Liquid density of the binary system $\{\text{CO}_2 + \text{H}_2\text{O}\}$ at 59 °C and 30 – 290 bar predicted by VTPR, T-VTPR vs. experimental data (Li et al., 2004).

4.2.3 Water binary systems with CO₂, H₂S, CH₄: CPA-PR, (T-)VTPR, TPR, GERG-2008

The comparative study of different EOS on the liquid density of water binary systems at high TP conditions was carried out, considering the CPA-PR model (Hajiw et al., 2015) as a reference. Fig. 4.9 shows the molar volume and the corresponding excess molar volume for three binaries {CH₄+H₂O}, {CO₂+H₂O}, {H₂S+H₂O} at 100 °C and 300 bar given by TPR, VTPR, T-VTPR, CPA-PR, GERG-2008 (REFPROP). The solubility of each component at the given TP was calculated by CPA-PR, then varying the component mole fraction, the liquid density was calculated up to the saturation point. Table 4.11 presents the comparison of the liquid density, the error is given relatively to CPA-PR results.

Table 4.11 – Comparison of CPA-PR, VTPR, T-VTPR and GERG-2008. The error, relative to CPA-PR of the liquid density of the binary systems {CH₄+H₂O}, {CO₂+H₂O}, {H₂S+H₂O} at 100 °C and 300 bar, in %.

	GERG-2008	TPR	VTPR (TPR)	T-VTPR (TPR)
{CH ₄ +H ₂ O}	1.1	17.9	3.9	3.0
{CO ₂ +H ₂ O}	0.8	16.7	3.6	2.8
{H ₂ S+H ₂ O}	1.3	17.4	3.9	3.6

The molar volume of CPA is located between VTPR and T-VTPR with the deviation of 4% and 3%, respectively, Fig. 4.9. TPR provides a higher v_m of 21% deviation and consequently a lower density of 17% deviation that is about of the same degree as AAD for water density given by the PR78, CPR, GPR and HPR (the α modified PR EOS), compare with Table 4.2. The TPR and volume translation EOS, based on the PR78 with the same mixing rules and binary coefficients, yield identical excess molar volume v_m^E , less than that of CPA-PR for all three binaries, see Fig. 4.9. Because, by definition, v_m^E shows the deviation of the real mixture molar volume from the molar volume of the ideal mixture, the molecules of the real mixture require less volume than those of the ideal mixture for all three binaries. The CPA-PR molar volume is closer to that of the ideal mixture, than the results of VTPR models. The v_m^E variation is about 5×10^{-8} for CH₄ and CO₂ mixtures

and slightly larger for H₂S binary.

The mass liquid density is displayed in Fig. 4.10. The CH₄ binary density decrease with the growing CH₄ mole fraction due to its molar weight, which is less than that of H₂O. For the CO₂ binary, the density grows toward the CO₂ saturation except the GERG-2008 results. The H₂S binary density declines by using the CPA-PR, GERG-2008 and VTPR, their v_m slopes are greater than those of the T-VTPR and TPR, Fig. 4.9. In spite of 4% underestimation, the **VTPR** model can qualitatively predict the liquid density of the water binary systems with CH₄, CO₂, H₂S at high PT conditions. For the system CO₂ + H₂O, the **T-VTPR** produces a good estimation of the liquid density with an acceptable overestimation over a wide range of PT .

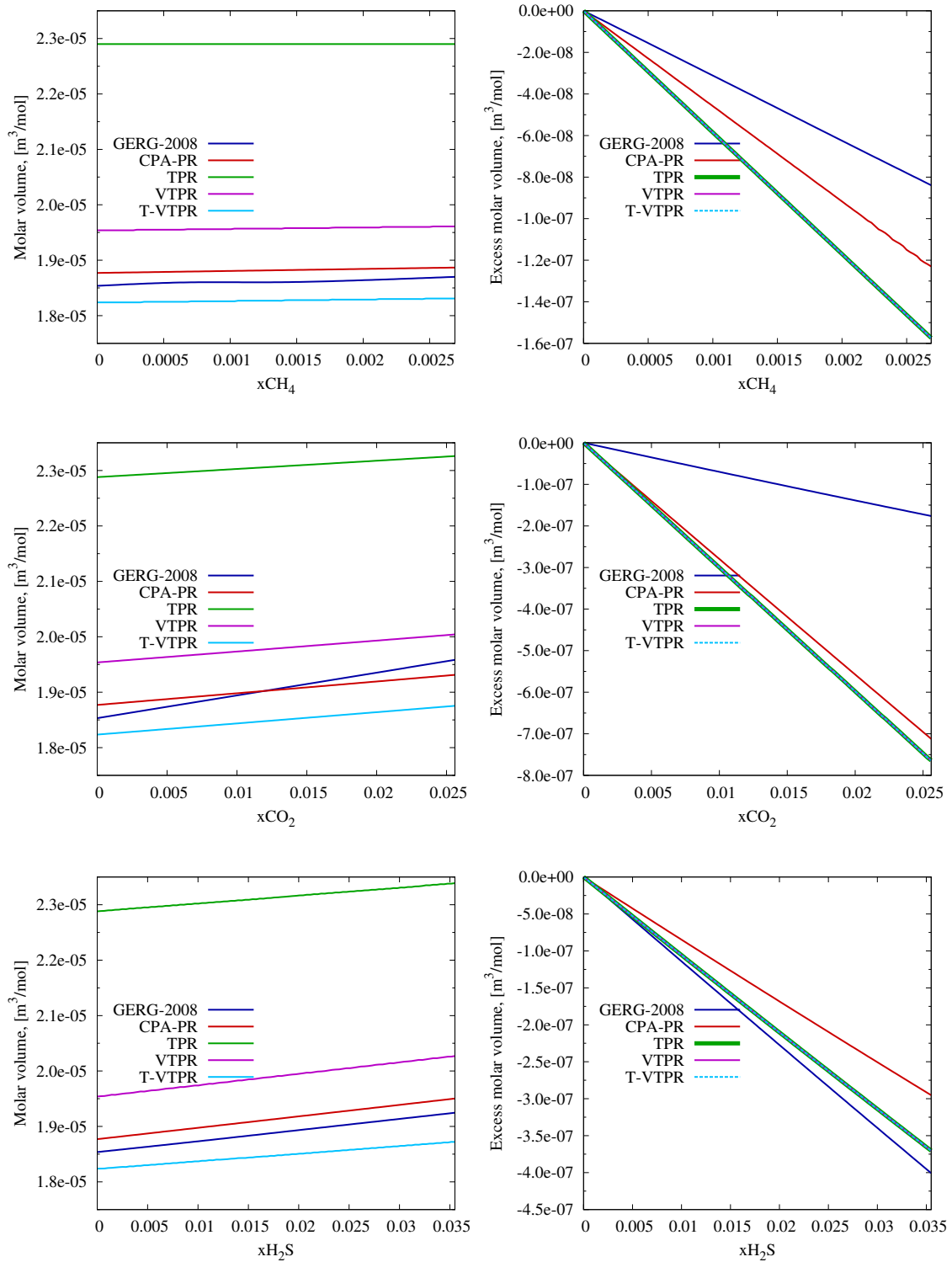


Figure 4.9 – Molar volume (left) and excess molar volume (right) of water binaries with CH_4 (top), CO_2 (center), H_2S (bottom) at 100 °C and 300 bar, predicted by GERG-2008 (REFPROP), CPA-PR, VTPR, T-VTPR.

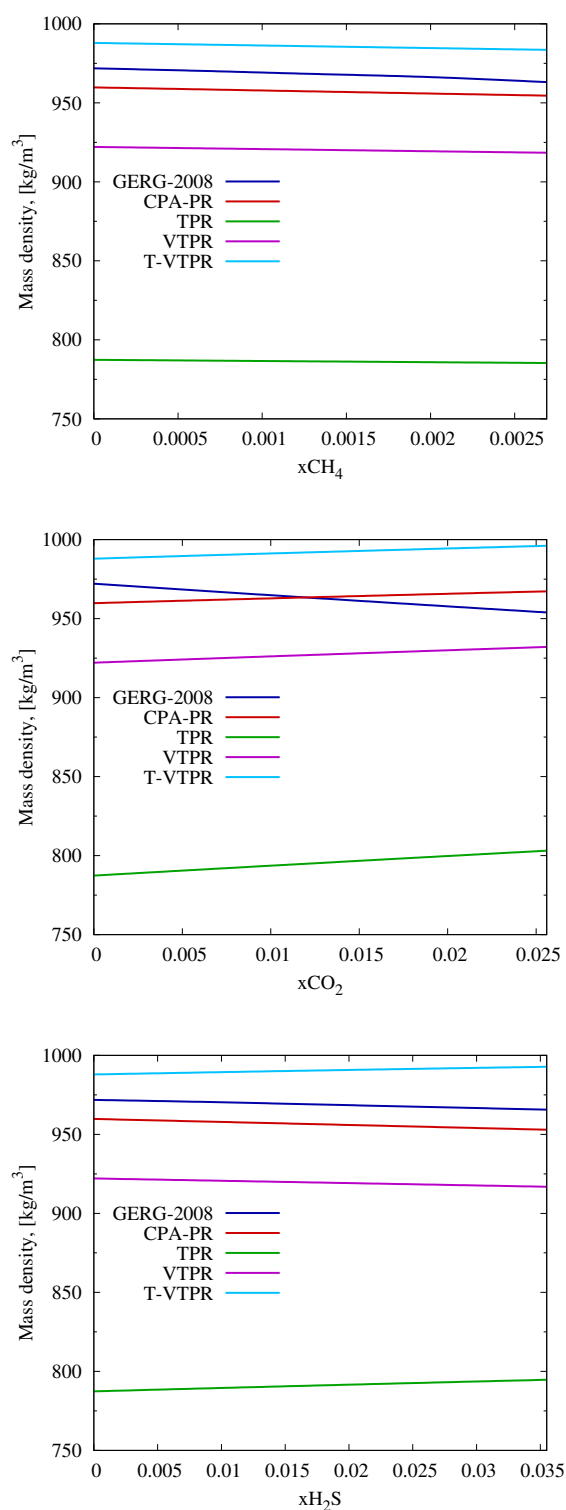


Figure 4.10 – Liquid density of water binaries with CH₄ (top), CO₂ (center), H₂S (bottom) at 100°C and 300 bar, predicted by GERG-2008 (REFPROP), CPA-PR, VTPR, T-VTPR.

4.3 Fluid properties

4.3.1 Density

Table 4.12 presents the available fluid density models in HYTEC, where the constant mode for both phases implies incompressible flow. The majority of density options stem from the EOS models, although the analytical formulation for brine/formation water is also possible by adjusting the measured fluid properties. The EOS description was given previously in Sec. 4.1.

Table 4.12 – List of the density models available in HYTEC.

Gas density model	Liquid density model
Constant	Constant
Ideal gas	Linear
PR78	PR78
PPR78	PPR78
TPR + (g)VTPR	TPR + (g)VTPR
TPR + (g)T-VTPR	TPR + (g)T-VTPR
	McCain

The linear and McCain models use the total dissolved solids (TDS). The former simply yields

$$\rho_l(S, T) = \rho_w(T)(1 + S), \quad (4.3.1)$$

where S is the TDS in mass fraction. Despite its trivial definition, applying this model gives a rough estimation of the phase transfer that arises from the solid-liquid and additionally from the liquid-gas interaction.

By assuming the primary NaCl content of formation water, McCain Jr. derived the liquid density $\rho_l(S, T, P)$ based on fluid samples data from the numerous petroleum reservoirs:

$$\rho_l(S, T, P) = \frac{\rho_w(S)}{B_w(P, T)}, \quad (4.3.2)$$

where ρ_w is the density of formation water at standard temperature pressure (STP) con-

ditions (25 °C, 1 atm), calculated by

$$\rho_w = 62.368 + 0.438603S + 1.60074 \times 10^{-3}S^2, \quad (4.3.3)$$

where S in wt% and B_w is the formation volume fraction of formation water, the temperature and pressure dependent function given by

$$B_w = (1 + \Delta V_{wP})(1 + \Delta V_{wT}), \quad (4.3.4)$$

$$\Delta V_{wP} = -1.95301 \times 10^{-9}PT - 1.72834 \times 10^{-13}P^2T \quad (4.3.5)$$

$$-3.58922 \times 10^{-7}P - 2.25341 \times 10^{-10}P^2, \quad (4.3.6)$$

$$\Delta V_{wT} = -1.0001 \times 10^{-2} + 1.33391 \times 10^{-4}T + 5.50654 \times 10^{-7}T^2, \quad (4.3.7)$$

where T and P are in °F and in psia, respectively. The validity of model was given with 2% deviation from the experiment data for $P < 350$ bar, $T < 127$ °C and $S < 30$ wt%, McCain Jr. (1991). Adams and Bachu (2002) displayed the reliability of McCain method providing the comparative study of five analytical density models for the water with the NaCl content up to 30 wt% at high TP conditions and for the formation waters of the Alberta Basin at $S < 27$ wt% at STP conditions; even though the underestimation of the measured densities increases with S , reaching about 2% at $S = 30$ wt%.

4.3.2 Viscosity

Modeling CO₂

Altunin (1975) developed the thermodynamic models for CO₂ and in particular, the viscosity model as a function of ρ and T for the range 220 – 1300 K and up to 1200 bar (Altunin, 1975, Eq. 9.33):

$$\mu_{CO_2} = \mu^0 \exp \left[\sum_{i=1}^4 \left(\sum_{j=0}^1 \frac{a_{ij}}{T_r^j} \right) \rho_r^i \right], \quad (4.3.8)$$

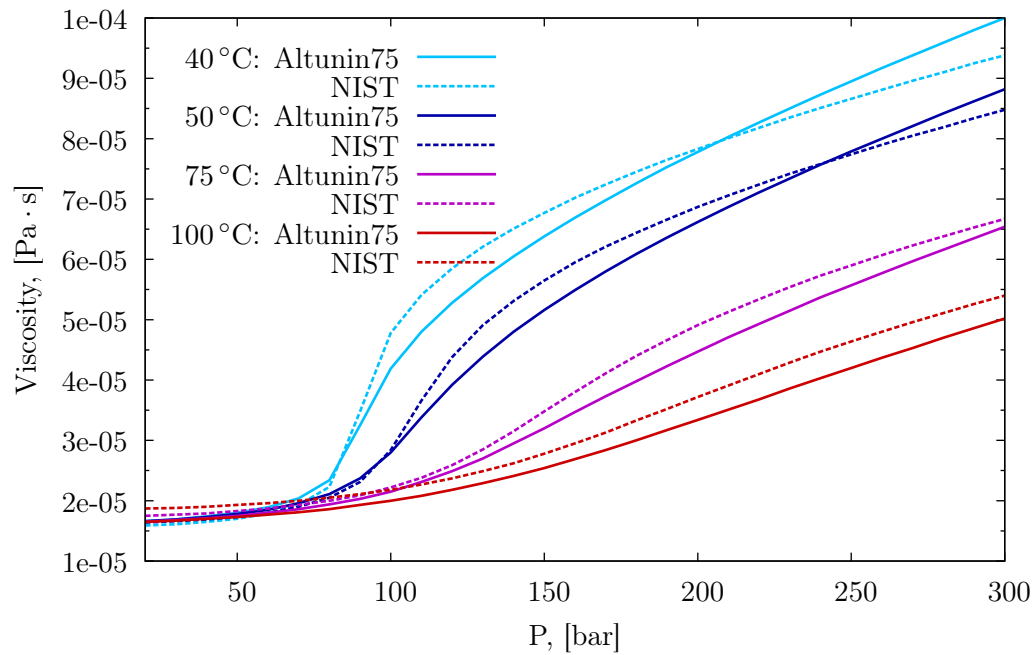
$$\mu^0 = \sqrt{T_r} \left(27.2246461 - \frac{16.6346068}{T_r} + \frac{4.66920556}{T_r^2} \right), \quad (4.3.9)$$

where $\rho_r = \rho/\rho_c$ and T_r are the reduced gas density and temperature, a_{ij} are the constant parameters given in Table 4.13, the density can be provided by EOS. The correlation of Altunin viscosity model with the database of Vargaftik (1975) presented in Garcia (2003) results in the relative deviation $< 2\%$ at $P > 150$ bar, while it reaches 8% at 50 – 70 °C

Table 4.13 – Parameters a_{ij} for the Altunin viscosity model of CO₂, Eq. (4.3.8).

a_{10}	$=$	0.248566120	a_{30}	$=$	0.3638545230
a_{11}	$=$	0.004894942	a_{31}	$=$	−0.7742290210
a_{20}	$=$	−0.373300660	a_{40}	$=$	−0.0639070755
a_{21}	$=$	1.227534880	a_{41}	$=$	0.1425070490

and 100 bar (for density, the Altunin model was applied, Altunin (1975)). Fig. 4.11 depicts the results of Altunin model by using the PR78 model for the density. The temperature and pressure dependance of the viscosity function follows that of the density, compared with Fig. 4.1.

**Figure 4.11** – Viscosity of CO₂ at 45, 50, 75 and 100 °C and 20 – 300 bar predicted by Altunin model Eq. (4.3.8) and NIST data.

Later, Fenghour et al. (1998) proposed the thermodynamic model for the viscosity of CO₂ as a function of density, T and P that is reliable under the high TP conditions with

the correction near the critical region, developed by Vesovic et al. (1990):

$$\mu_{\text{CO}_2}(\rho, T) = \mu_0(T) + \Delta\mu(\rho, T) + \Delta\mu_c(\rho, T), \quad (4.3.10)$$

where μ_0 is the viscosity in the zero-density limit, $\Delta\mu$ is the excess viscosity and $\Delta\mu_c$ is the critical enhancement. The first and the second terms consists of the polynomial forms of 4 on T and of 8 on P , respectively. The critical correction requires the matrix multiplication and the solution of cubic equation. Fenghour et al.'s model results in 2% of accuracy at $T < 260$ K and $P < 250$ MPa for the liquid viscosity. Since the comparison with Altunin formulation showed 2% of deviation in the gas and supercritical regions (Garcia, 2003), the latter can be chosen for the CO_2 viscosity modeling at corresponding TP conditions.

Modeling H_2O , $\{\text{H}_2\text{O} + \text{CO}_2\}$ and $\{\text{H}_2\text{O} + \text{NaCl} + \text{CO}_2\}$

The Islam and Carlson (2012) model describes the H_2O viscosity at the TP range 20 – 105 °C, 0 – 600 bar, Eq. (4.3.11), producing a maximum error 0.05 of the IAPWS data of 2009. Then, the authors proposed the formulation for $\{\text{H}_2\text{O} + \text{CO}_2\}$ system, Eqs. (4.3.12) and (4.3.13). The error up to 1% occurs in the TPx range 20 – 100 °C, 50 – 450 bar and 0 – 0.024 mole fraction.

$$\mu_{\text{H}_2\text{O}} = a_0 + \sum_{i=1}^3 b_i \exp(-c_i T) + P \sum_{i=0}^3 d_i (T - 293.15)^i, \quad (4.3.11)$$

$$\mu_{\{\text{H}_2\text{O}, \text{CO}_2\}} = \mu_{\text{H}_2\text{O}} \mu_r, \quad (4.3.12)$$

$$\mu_r = 1 + \frac{\sum_{i=1}^2 a_i^r x_{\text{CO}_2}^i}{\sum_{i=0}^1 b_i^r T^i}, \quad (4.3.13)$$

where a_i , b_i , c_i , d_i and a_i^r , b_i^r are the constant parameters, see Tables 4.14 and 4.15. The viscosity is strongly temperature dependent and slightly pressure dependent, the CO_2 content impact decreases with the temperature, see Fig. 4.12.

For the $\{\text{H}_2\text{O} + \text{NaCl} + \text{CO}_2\}$ viscosity, Islam and Carlson derived the following equation

$$\mu_{\{\text{H}_2\text{O}, \text{NaCl}, \text{CO}_2\}} = \mu_{\{\text{H}_2\text{O}, \text{NaCl}\}} \left(1 + 4.65 x_{\text{CO}_2}^{1.0134} \right), \quad (4.3.14)$$

where x_{CO_2} is the molality of CO_2 . The $\{\text{H}_2\text{O} + \text{NaCl}\}$ viscosity formulation will be presented below, Eq. (4.3.15).

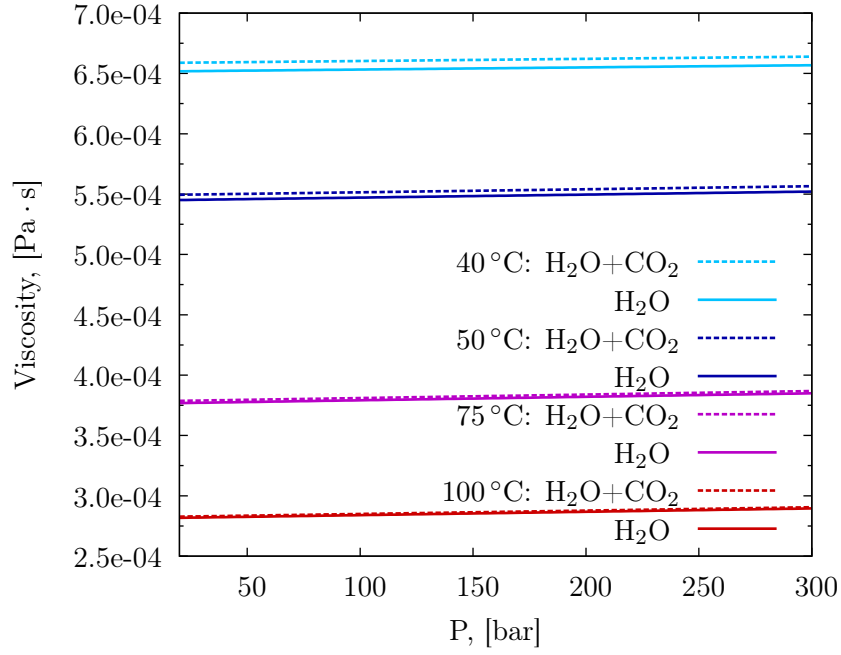


Figure 4.12 – Viscosity of H_2O and $\text{H}_2\text{O} + \text{CO}_2$ with $x_{\text{CO}_2} = 0.02$ at 45, 50, 75 and 100 °C and 20 – 300 bar predicted by Islam-Carlson models Eqs. (4.3.11) and (4.3.12).

Modeling $\{\text{H}_2\text{O} + \text{NaCl}\}$

The Mao and Duan (2009) model predicts the viscosity with the accuracy about 1% for binary systems including $\{\text{H}_2\text{O} + \text{NaCl}\}$ up to 6 mol/kg at the TP range 273 – 623 K and 1 – 1000 bar:

$$\mu_{\{\text{H}_2\text{O}, \text{NaCl}\}} = \mu_{\text{H}_2\text{O}} \mu_r^{MD}, \quad (4.3.15)$$

$$\ln \mu_r^{MD} = Am + Bm^2 + Cm^3, \quad (4.3.16)$$

where μ_r^{MD} is the relative viscosity, m is the molality of salts, A , B , C are the temperature dependent functions:

$$A = \sum_{i=0}^2 A_i T^2, \quad (4.3.17)$$

$$B = \sum_{i=0}^2 B_i T^2, \quad (4.3.18)$$

$$C = \sum_{i=0}^1 C_i T^2. \quad (4.3.19)$$

Table 4.14 – Parameters a_i , b_i , c_i , d_i for the Islam and Carlson viscosity model of H_2O , Eq. (4.3.11).

$$\begin{array}{rcl}
 a_0 & = & 9.03591045 \times 10^1 \\
 b_1 & = & 3.40285740 \times 10^4 \\
 b_2 & = & 8.23556123 \times 10^8 \\
 b_3 & = & -9.28022905 \times 10^8 \\
 c_1 & = & 1.40090092 \times 10^{-2} \\
 c_2 & = & 4.86126399 \times 10^{-2} \\
 c_3 & = & 5.26696663 \times 10^{-2} \\
 d_0 & = & -1.22757462 \times 10^{-1} \\
 d_1 & = & 2.15995021 \times 10^{-2} \\
 d_2 & = & -3.65253919 \times 10^{-4} \\
 d_3 & = & 1.97270835 \times 10^{-6}
 \end{array}$$

Table 4.15 – Parameters a_i^r , b_i^r for the Islam and Carlson viscosity model of $\{\text{H}_2\text{O}+\text{CO}_2\}$, Eq. (4.3.13).

$$\begin{array}{rcl}
 a_1^r & = & 7.632609119 \times 10^2 \\
 a_2^r & = & -9.46077673946 \times 10^3 \\
 b_0^r & = & -1.047187396332 \times 10^4 \\
 b_1^r & = & 3.68325597 \times 10^1
 \end{array}$$

The constant parameters A_i , B_i , C_i for $\{\text{H}_2\text{O}, \text{NaCl}\}$ system are given in Table 4.16.

Fig. 4.13 displays the example of the HYTEC syntax used for the definition of the density, viscosity, relative permeability and capillary pressure models.

Table 4.16 – Parameters A_i , B_i , C_i for the Mao and Duan viscosity model of $\{\text{H}_2\text{O} + \text{NaCl}\}$, Eq. (4.3.15).

A_0	$=$	-0.21319213	B_1	$=$	$-0.27292263 \times 10^{-3}$
A_1	$=$	$0.13651589 \times 10^{-2}$	B_2	$=$	$0.20852448 \times 10^{-6}$
A_2	$=$	$-0.12191756 \times 10^{-5}$	C_0	$=$	$-0.25988855 \times 10^{-2}$
B_0	$=$	$0.69161945 \times 10^{-1}$	C_1	$=$	$0.77989227 \times 10^{-5}$

```

capillary-pressure = van-genuchten {
  entry-pressure = 19.61 kPa
  pow = 0.457
  slr = 0.0
  sgr = 0.0
}
liquid-relative-permeability-model = van-genuchten {
  pow = 0.457
  slr = 0.3
  sgr = 0.0
}
gas-relative-permeability-model = brooks-corey {
  pow = 2
  slr = 0.3
  sgr = 0.05
}
liquid-density-model = t-vtpr
gas-density-model = pr
liquid-viscosity-model = islam-carlson
gas-viscosity-model = altunin

```

Figure 4.13 – Example of the model definition of the fluid and rock-fluid properties in the HYTEC input file.

Chapter 5

Applications: CCS, Impact of Co-injected Impurities and Convective Mixing

Le Chapitre 5 présente des simulations numériques de stockage du carbone et s'est focalisé sur l'impact des impuretés co-injectées et le mélange convectif à l'échelle du laboratoire et du réservoir. La chaîne des simulations commence par le problème 1D de la séparation chromatographique qui a pour objectif de modéliser des phénomènes observés à l'échelle du laboratoire. Après, le problème convectif avec une couche de gaz fine est traité. Les résultats sont comparés à ceux de simulations numériques d'autres codes et d'estimations analytiques. Nous continuons avec la modélisation 2D radiale de stockage de CO_2 afin de mettre en évidence l'impact du flux convectif sur la dissolution. En outre, une formulation composée de deux modèles analytiques est utilisée pour évaluer la sous-estimation numérique de dissolution due à la convection. Le problème est ensuite étendu à l'injection de gaz complexes : trois scénarios d'injection montrent une forte interaction entre le flux convectif dense et la séparation chromatographique et leurs effets sur la vitesse du front gazeux et sur la dissolution globale de gaz.

This chapter presents the numerical and analytical study of carbon capture and storage (CCS) associated with the presence of gas impurities. During the capture process, CO_2 is separated to increase the efficiency of the storage and to respect the environmental and

legal aspects. Nevertheless, impurities such as N_2 , O_2 , Ar, SO_2 , CH_4 , H_2S can remain, (Kather, 2009). The thermodynamic properties of these CO_2 binary mixtures were studied experimentally and numerically by Al-Siyabi (2013); Ziabakhsh-Ganji and Kooi (2012); Li and Yan (2009); Ziabakhsh-Ganji and Kooi (2014). The authors highlighted the positive density and solubility effects of SO_2 and the negative impact of Ar, O_2 and N_2 on the volume and solubility storage capacity. In order to reveal the impurity impact on the reservoir scale, we study its behavior starting from a one-dimensional problem of the gas dynamics in a closed box. Next, we raise the issue of convective mixing in the aqueous phase in the contact with the gas and the capillary fringe, that is demonstrated on a 2D problem. Then, we compare the numerical results of a 2D radial CO_2 injection with some analytical solutions. Finally, the model is used to investigate the effect of the presence of impurities in the injected stream.

5.1 Methods

The numerical method described in Sec. 3.2 is employed to model the two-phase flow. The density properties of gas and liquid mixtures are predicted by the PR78 and T-VTPR models, respectively; we chose the binary interaction parameters given by Li and Yan (2009), that were calibrated for calculation VLE of CO_2 with the impurities. The preliminary work on the liquid density was carried out: the comparison of PR78, T-VTPR, CPA models and GERG-2008 (REFPROP) for pure components and binary systems involved in CCS, Sec. 4.2. In Al-Siyabi (2013), the PR78 was studied on eight CO_2 binary systems including O_2 , N_2 , SO_2 , during which the PR78 ADD for these three systems on the saturation pressure and on the vapor composition was evaluated as 3.2% and 1.9%, respectively. These EOS models were fitted to the experimental data acquired on binary systems. They can be applied on ternary or more systems, even though their accuracy could then be questioned, and would require datasets, not yet existing.

5.2 Gas dynamics and chromatographic partitioning

The chromatographic partitioning of injected gas was investigated experimentally in the work of Bachu and Bennion, in which the authors provided the sensitivity study on the impact of gas impurity concentration (2, 5, 30% of H_2S), PTS conditions (61°C and 13.5 MPa; 25°C and 6 MPa; 119 ppm) and of admixture type in the injected stream (H_2S , N_2 , SO_2 , CH_4). In the last stage, the laboratory results revealed a key factor of the impurity solubility order regarding to that of CO_2 . The breakthrough time of the gas compounds and their profiles highly depend on the solubility ratio between gases. In Fig. 5.1, the mole fraction of the impurity gas H_2S , N_2 , SO_2 and C_1 (methane) are normalized by the relevant initial mole fraction of the injected gas (5% for all four tests) in order to emphasize the scale of the gas partitioning effect due to the different solubility relations. The less soluble the admixture gas is, the earlier and higher breakthrough is expected. The impurity gases present in the injection stream after the capture processes can then be classified by the solubility. For example, we can write them in the decreasing order of the solubility (Henry's

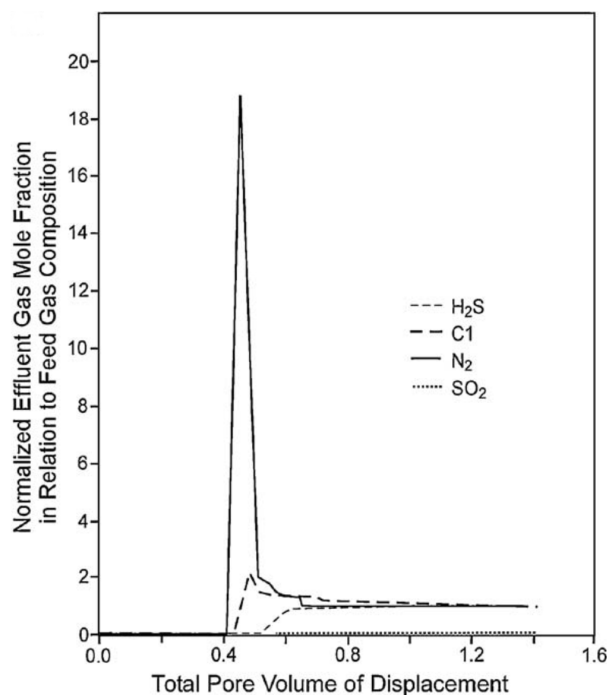


Figure 5.1 – Mole fraction of H_2S , C_1 (methane), N_2 and SO_2 at the outlet normalized by the relevant mole fraction in the initially injected gas, as a function of the pore volume of the total injection (Bachu and Bennion, 2009).

constant in \log) at 60°C :

$$\begin{aligned} \text{SO}_2(0.221) &> \text{H}_2\text{S}(1.265) > \text{CO}_2(1.767) \\ &> \text{Ar}(3.016) > \text{CH}_4(3.023) > \text{O}_2(3.063) > \text{N}_2(3.325). \end{aligned} \quad (5.2.1)$$

Hence, the gases on the RHS to CO_2 of the inequality (from Ar to N_2) should evolve the similar profile of the normalized mole fraction with high and early peaks at the front, and vice versa, SO_2 and H_2S yield the lower and later breakthrough. Bachu and Bennion (2009) demonstrated experimentally this effect in the case of H_2S , N_2 , SO_2 , CH_4 and later, numerically for H_2S admixture Bachu et al. (2009). To complete sequence (5.2.1), we propose the simulation of the following problem where the resident gas is represented by pure CO_2 and then by $\text{CO}_2 + 5\%$ of impurities: SO_2 , Ar, O_2 , N_2 .

5.2.1 Problem description

The homogeneous medium is 10 cm long, and 1 cm high. Its right half is fully saturated by water. The left one contains some amount of gas ($S_g = 0.2$), the water is at equilibrium with the gas phase. To exclude the gravity forces, the geometry is simplified to 1D, also the liquid density is supposed to be constant. Supposing the PT conditions of a deep reservoir, the temperature is 50°C and the initial pressure is 100 bar. The parameters of the problem are given in Table 5.1. We firstly present the results for the case when the gas consists only of CO_2 . Then, the impact of impurities is observed on the example of SO_2 and N_2 , as they are the most and the least soluble gases of sequence 5.2.1. Finally, all five systems are compared: CO_2 , $+\text{SO}_2$, $+\text{Ar}$, $+\text{O}_2$, $+\text{N}_2$.

5.2.2 CO_2

Driven by the capillary forces, the gas, CO_2 in this case, moves forward from the left to the right, regressively occupying the porous medium and dissolving in the fresh water until

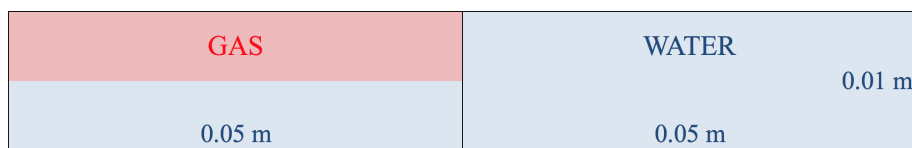


Figure 5.2 – Gas dynamics and chromatographic partitioning: geometry of the problem.

Table 5.1 – Parameters of the 1D problem of the chromatographic partitioning.

Initial Conditions		
	$\Omega_g = [0, 0.05] \times [0, 0.01] \text{ [m}^2\text{]}$	$\Omega_w = [0.05, 0.1] \times [0, 0.01] \text{ [m}^2\text{]}$
	$p_{l0} = 100 \text{ [bar]}, S_{g0} = 0.2$	$p_{l0} = 100 \text{ [bar]}, S_{g0} = 0$
Temperature	50 °C	
Boundary Conditions		
$\partial\Omega :$	no flow	
Matrix Properties		
Porosity ϕ	0.2	
Permeability K	$10^{-12} \text{ [m}^2\text{]}$	
Fluid-Matrix Properties		
Brooks-Corey capillary pressure and relative permeability		
Eq. 2.1.7, Eq. (2.1.9), Eq. (2.1.10): $p_b = 10^5 \text{ Pa}$, $\lambda = 2$, $S_{lr} = 0.15$, $S_{gr} = 0.05$		
Fluid Properties		
Density	$\rho_l = 1008.19$	ρ_g : PR78
Viscosity	$\mu_l = 5.49 \times 10^{-5}$	$\mu_g = 2.84 \times 10^{-5} \text{ [Pa} \cdot \text{s]}$
Diffusion coefficient	$D_l = 3.36 \times 10^{-9}$	$D_g = 1.95 \times 10^{-7} \text{ [m}^2\text{/s]}$

reaching its stationary state, Fig. 5.3 (left). During the continuous gas dissolution, both pressures, liquid and gas, decrease. In Fig. 5.3 (right), the temporal evolution of the phase pressures is plotted at $x = 0.005$, at the left border of the domain. The shape of the pressure function is correlated with the decreasing dissolution rate as the gas approaches slower to the right border.

5.2.3 CO₂ + 5% impurities

Supposing that the gas is represented by 95 mol% of CO₂ and 5 mol% of impurities, we compare the numerical results for N₂ and SO₂ (SO₂ is more soluble than CO₂ and N₂ is less soluble than CO₂). The chromatographic partitioning of gas composition occurs arising from the different solubility rates, as expected, Fig. 5.4. The peak of the N₂ mole fraction is 0.97 at 10 s, so the normalized value is 19.4 that corresponds to the observations

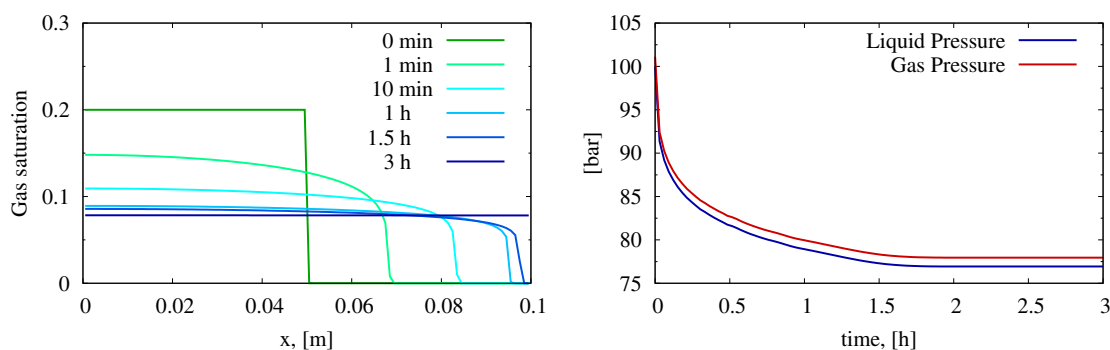


Figure 5.3 – Temporal evolution of gas saturation (left), the liquid and gas pressures in bar (right) at point $x = 0.005$.

of Bachu and Bennion, Fig. 5.1. The N_2 mole fraction decreases with time, afterwards. Nevertheless the mole fraction normalized to its initial value (5 mol%) is higher during the simulation time, until the stationary state is reached. The gas partitioning of CO_2 and SO_2 is opposite to the system $\{CO_2 + N_2\}$, CO_2 takes a leading position of the gas front, Fig. 5.4 (right column).

The partitioning of the aqueous components can be found in addition to that of the gas phase. Being less soluble than CO_2 , N_2 forms a slight accumulation ahead towards the displacement, which rapidly diminishes. Meanwhile, the high solubility of SO_2 yields the pronounced concentration behind the moving front. Fig. 5.5 shows the concentration contours of aqueous components for both systems.

Bachu and Bennion reported the unexpected earlier breakthrough of CO_2 in the cases of N_2 and CH_4 suggesting that these gases pull CO_2 along. Based on the present results, we confirm the associated movement of CO_2 along with the impurities. Fig. 5.6 (right column) demonstrates the advancing front of gas saturation of the system with less soluble N_2 and consequently, a delay of gas saturation profile of the system $CO_2 + SO_2$. The different position of the fronts is also conditioned by the gas density. Fig. 5.6 (left column) illustrates that the admixtures result in a high deviation of gas density: SO_2 increases the density, whereas N_2 reduces it. The gas chromatographic partitioning implies a density heterogeneity. In this problem, such important changes in gas mixture density determine the dynamics of the system and, in particular, for the gas saturation front. The models of CO_2 and $\{CO_2 + N_2\}$ achieve the equilibrium state within one day, while it takes about

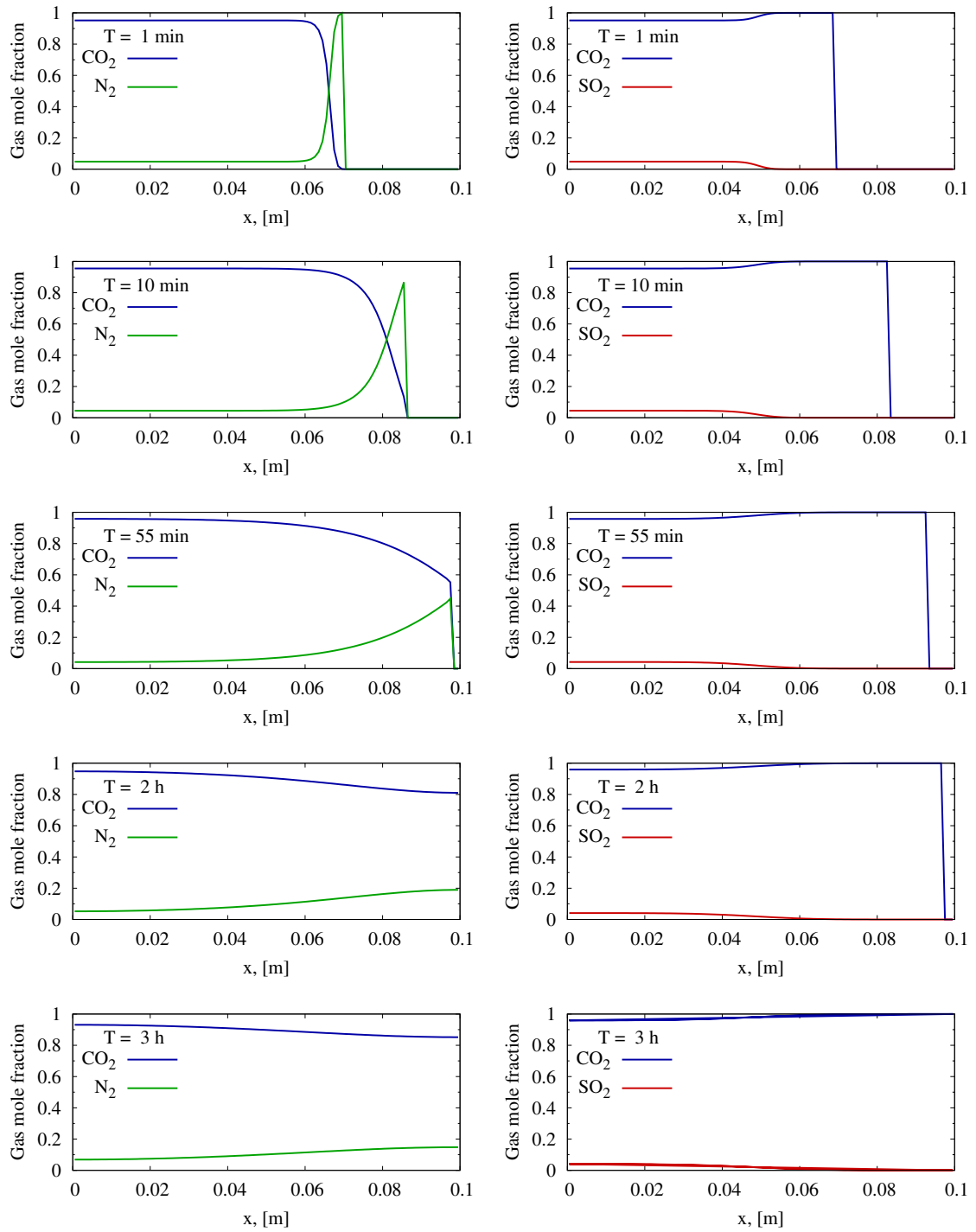


Figure 5.4 – Gas composition of $\{\text{CO}_2 + \text{N}_2\}$ (left) and $\{\text{CO}_2 + \text{SO}_2\}$ (right) in mole fraction at $T = 1$ min (top), 10 min, 55 min, 2 h and 3 h (bottom).

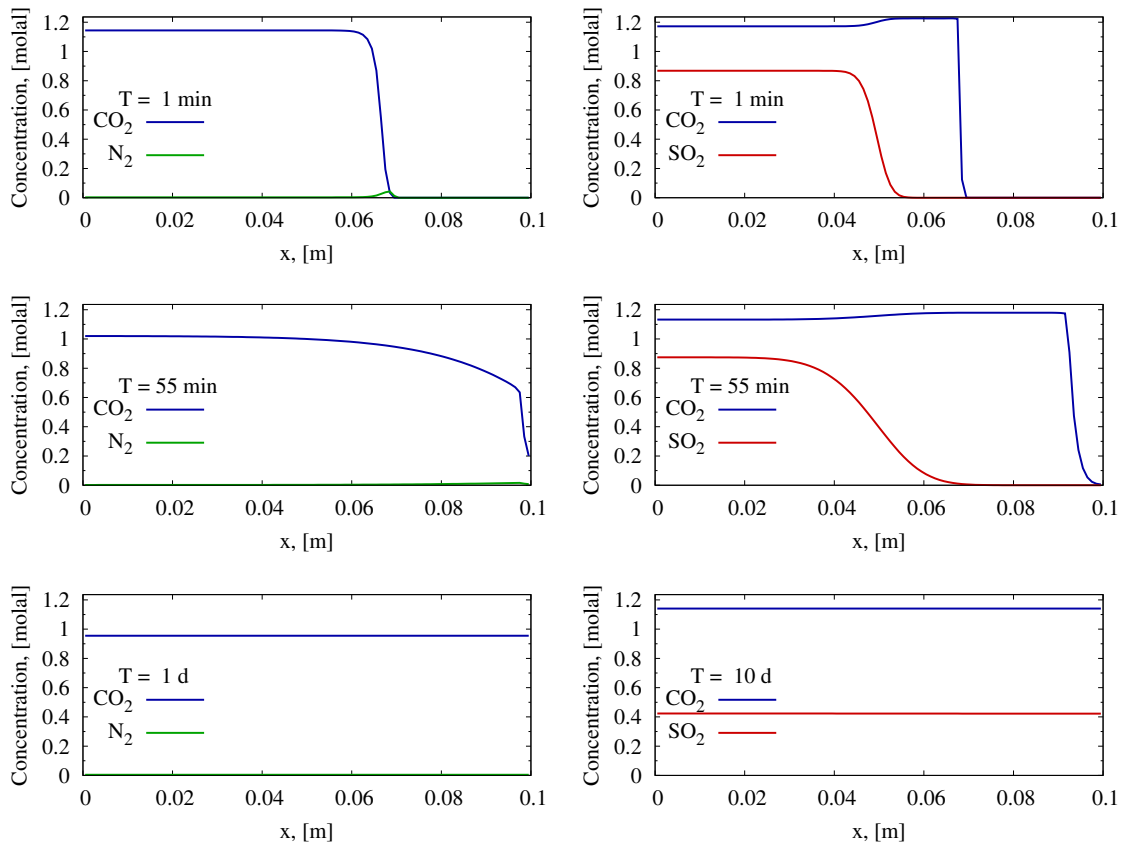


Figure 5.5 – Aqueous composition of $\{CO_2 + N_2\}$ (left) and $\{CO_2 + SO_2\}$ (right) in molal at $T = 1$ min (top), 55 min (center), 1 and 10 d (bottom). The scale of the figures is identical, the concentration range lies in $[0; 1.24]$.

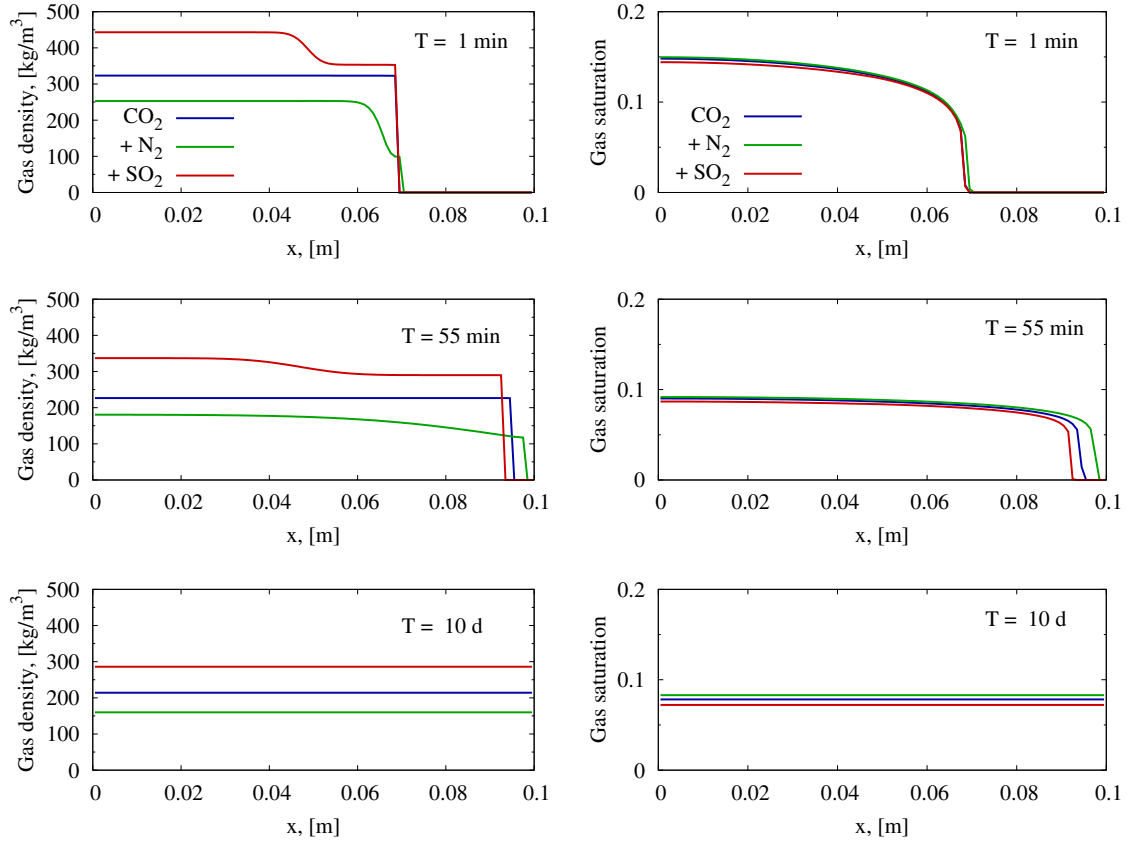


Figure 5.6 – Gas density in kg/m^3 (left) and saturation (right) of CO_2 , $\{\text{CO}_2 + \text{N}_2\}$ and $\{\text{CO}_2 + \text{SO}_2\}$ models at $T = 1$ min (top), 55 min (center), 1 and 10 d (bottom).

10 days in the $\{\text{CO}_2 + \text{SO}_2\}$ case.

The group of less soluble gases, Ar, O_2 , N_2 , provides similar results, Fig. 5.7. The gas saturation profiles of Ar and O_2 are very close due to their almost equal solubilities, see (5.2.1). The ordering of the gas admixture mole fractions at the stationary state corresponds inversely to that of the solubility:

$$\text{SO}_2(0.019) < \text{Ar}(0.097) < \text{O}_2(0.099) < \text{N}_2(0.104). \quad (5.2.2)$$

The CO_2 concentration is lower in the presence of Ar, O_2 , N_2 than that of the pure CO_2 and SO_2 impurity cases, Figs. 5.7 and 5.8. At equilibrium state, the concentration of CO_2 is the highest in the $\{\text{CO}_2 + \text{SO}_2\}$ system. However, that does not involve a better mass solubility capacity, as the initial volume is identical for the models but not the initial mass of CO_2 .

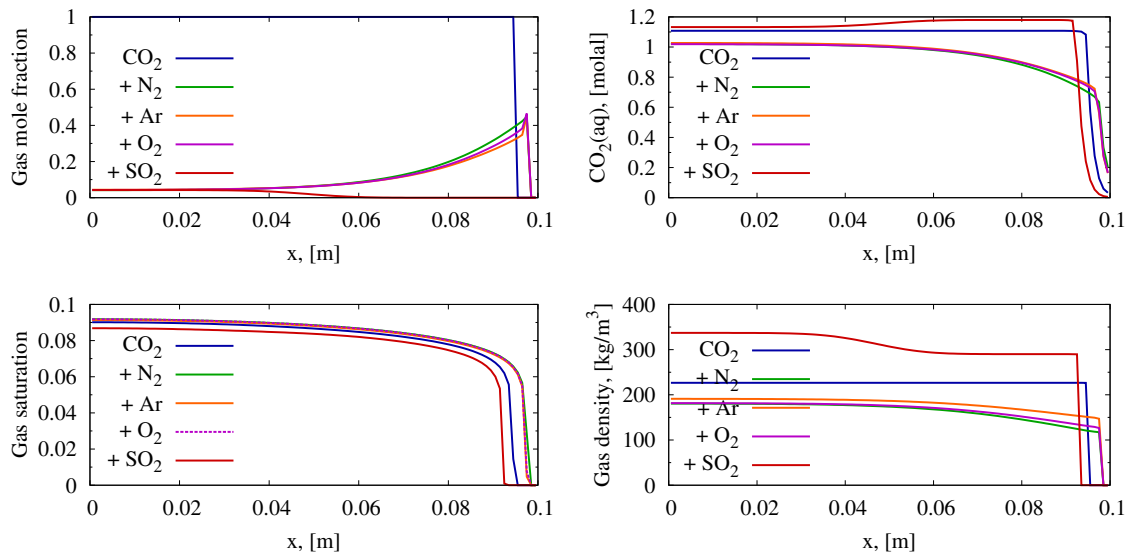


Figure 5.7 – Gas mole fractions of impurity gases and pure CO_2 (left top), CO_2 concentration in molal (right top), gas saturation (left bottom) and the gas density in kg/m^3 (right bottom) of CO_2 , $\{\text{CO}_2 + \text{N}_2\}$, $\{\text{CO}_2 + \text{Ar}\}$, $\{\text{CO}_2 + \text{O}_2\}$ and $\{\text{CO}_2 + \text{SO}_2\}$ models at $T = 55$ min.

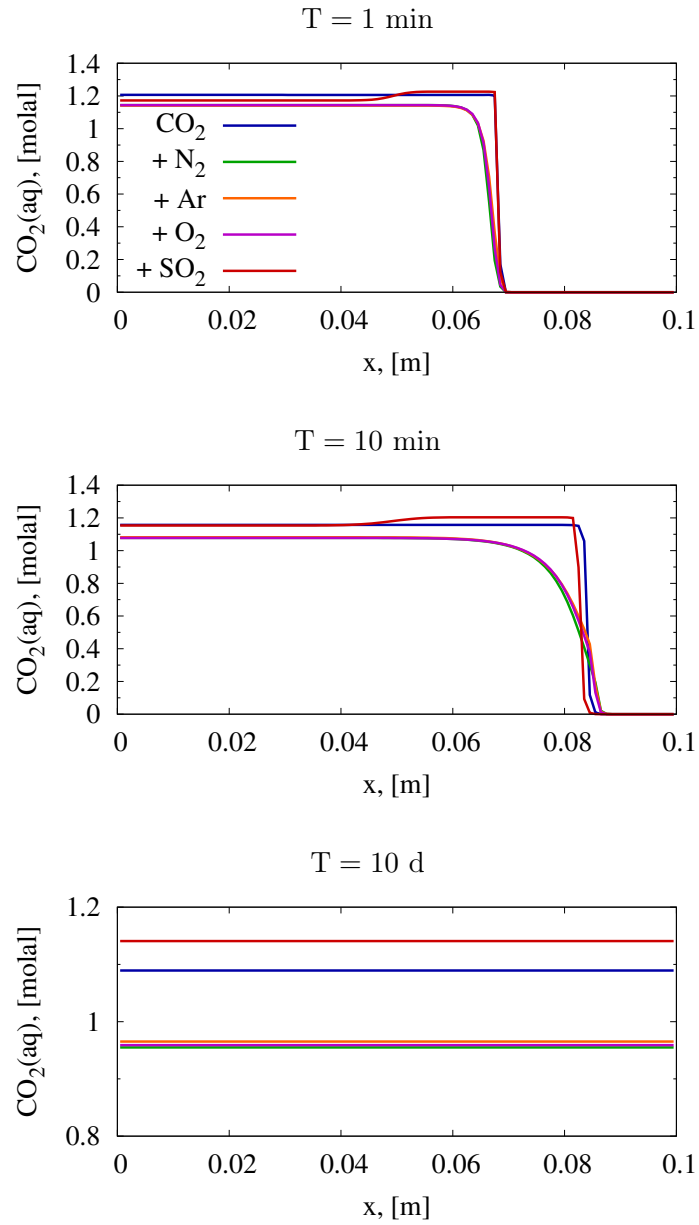


Figure 5.8 – CO_2 concentration in molal of CO_2 , $\{\text{CO}_2 + \text{N}_2\}$, $\{\text{CO}_2 + \text{Ar}\}$, $\{\text{CO}_2 + \text{O}_2\}$ and $\{\text{CO}_2 + \text{SO}_2\}$ models at $T = 1 \text{ min}$ (top), 10 min (center) and 10 d (bottom).

5.3 Convective mixing

In the context of CO_2 geological sequestration, the dissolution is identified as an important long-term trapping mechanism. As the supercritical CO_2 is buoyant to the ambient brine, the current spreads along the reservoir or aquifer inducing a security issue of the CO_2 leakage. The enhanced dissolution by convective mixing is therefore a favorable aspect of carbon storage. The formation water containing the dissolved CO_2 becomes heavier with increasing amount of dissolved CO_2 . The injected gas migrates, then accumulates at the top of the geological structure creating the $\text{CO}_2(\text{aq})$ saturated current denser than the formation water. The contact zone becomes unstable and the fingers enriched with dissolved CO_2 grow and sink down, initiating a convective flux and consequently, a reverse upward motion of the ambient brine, Fig. 5.9. We recall that the gases such as Ar, CO_2 and SO_2 increase the mass liquid density of water system, while adding CH_4 , N_2 , O_2 and H_2S makes it lighter.

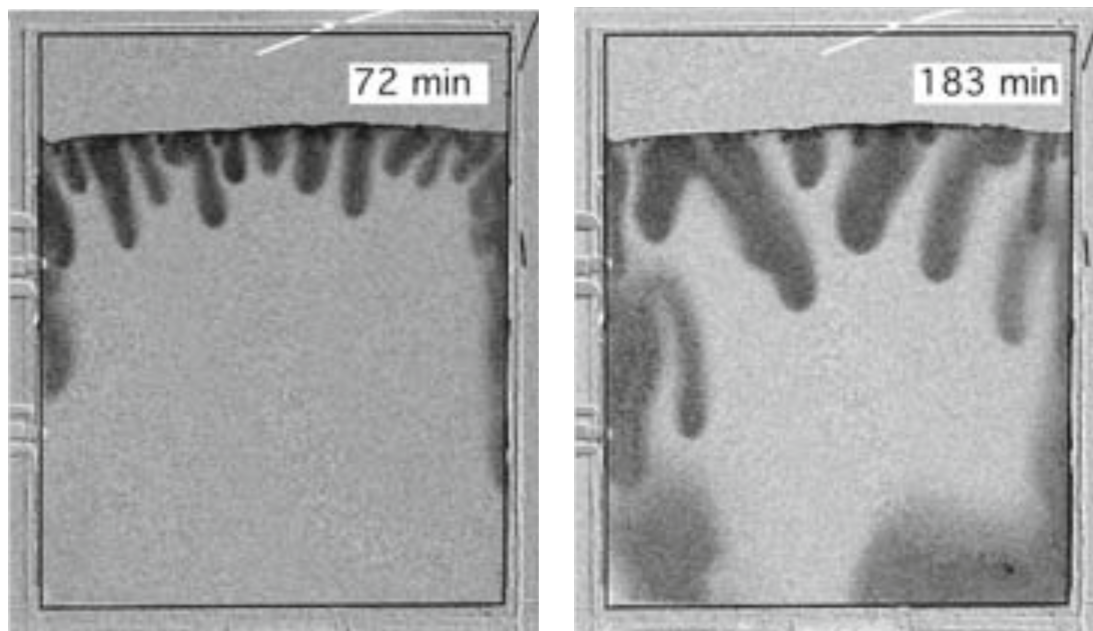


Figure 5.9 – Density driven convection at 72 (left) and 183 min (right), Kneafsey and Pruess (2010). Hele-Shaw cell ($25.4 \text{ cm} \times 30.5 \text{ cm}$) with the open top contains pure water, the injection of $\text{CO}_2(\text{g})$ is performed by a thin tube (white line at the top) at a distance from the gas/water interface. Higher $\text{CO}_2(\text{aq})$ concentration is represented by darker grey.

Lindeberg and Wessel-Berg (1997) derived a criterion of the stability for the convective motion based on the reservoir parameters $Ra < 4\pi^2$,

$$Ra = \frac{KgH\Delta\rho}{\mu\phi D}, \quad (5.3.1)$$

where H is the height, $\Delta\rho$ is the density increase due to the dissolved CO_2 . In Eq. 5.3.1, Ra is a concentration part of the S-number that Lindeberg and Wessel-Berg defined for the thermal and concentration convection criterion. Taking the typical North Sea reservoir parameters (Lindeberg and Wessel-Berg, 1997), one can estimate Ra of order 3 or 4, indicating the positive reservoir conditions for the convective mixing.

Farajzadeh et al. (2007), Kneafsey and Pruess (2010), Neufeld et al. (2010), MacMinn et al. (2012) showed experimentally the convective flow and its importance as a natural enhancement of CO_2 dissolution. Neufeld et al. (2010) and MacMinn et al. (2012) used a mixture of methanol and ethylene glycol to imitate the density properties of CO_2 ; also these authors unveiled the analytical formulation of the current spreading (MacMinn et al., 2012) and the convective flux for the stationary layer neglecting the diffusion and dispersion impact on the dissolution (Neufeld et al., 2010):

$$F_{\text{CO}_2} = \frac{\phi DX_{\text{CO}_2}}{H} Sh, \quad (5.3.2)$$

$$Sh = \alpha Ra^{4/5}, \quad (5.3.3)$$

where X_{CO_2} is the mass fraction of $\text{CO}_2(\text{aq})$, Sh is the Sherwood number, $\alpha = 0.19 \pm 0.01$. Applying linear stability analysis to the convective problem in the single phase only with diffusion, Riaz et al. (2006) derived the wavelength λ_c and the onset time of instability t_c . Then Elenius et al. (2012, 2014) considered a capillary transition zone that introduced the vertical flux at the instability interface and accelerated the convection:

$$\lambda_c \in \left[\frac{2\pi\mu D}{0.086K\Delta\rho g}, \frac{2\pi\mu D}{0.07K\Delta\rho g} \right], \quad (5.3.4)$$

where the left limit includes the capillary forces, the right limit is given for the diffusive dominant mode, Riaz et al. (2006). The onset time is expressed as follows

$$t_c = c_0 D \left(\frac{\phi\mu}{K\Delta\rho g} \right)^2, \quad (5.3.5)$$

where c_0 is the constant:

- Riaz et al. (2006) derived $c_0 = 146/\phi$, about the order of 2 or 3;

- Xu et al. (2006) proposes $c_0 = 75.19$;
- Elenius et al. (2012) suggests $c_0 = 31$ taking the capillary transition zone into account;
- Pau et al. (2010) – $c_0 = 1796$ and 3670 for porosity and permeability fluctuations;
- Li and Jiang (2014) – $c_0 = 1231$.

In the presence of the gas layer, the capillary forces create additional perturbations. The development of the density driven fingering can entail the uneven contact surface between the gas and liquid phases. The subject is raised in this section by 2D numerical modeling.

5.3.1 Problem description

We model the convective mixing problem with a thin $\text{CO}_2(\text{g})$ layer at the top of the 2D rectangle domain supplying $\text{CO}_2(\text{aq})$ at the upper boundary by pure diffusion process. Table 5.2 gives the problem parameters. The fluid, rock properties are identical to those of the previous modeling, Table 5.1. The liquid density is calculated using the T-VTPR model whose prediction of the $\text{CO}_2 + \text{H}_2\text{O}$ density and, especially, its increase are acceptable, Sec. 4.2.2. Given these fluid/matrix parameters, $Ra = 9728$, so the convective flow occurs. The wavelength estimation by Eq. 5.3.4 yields $\lambda_c \in [0.15, 0.184]$ m, the onset time lies in the range $t_c \in [0.45, 26.13]$ d. The width W of the domain should be greater than the wavelength, $W/\lambda_c = 54.2$ in this problem. The cell length is required to be smaller than the wavelength in order to capture the density fingering development. In the domain of interest (where the fingers are formed), the grid is uniform. We model the chain of simulations with $h_{x,z} = 1/16, 1/8, 1/4, 2$ [m]. For the finest grid, $\lambda_c/h_{x,z} = 2.95$, $H = 20$ m at a depth of $z_D = 1020$ m, the grid dimension is 51,200 cells.

5.3.2 Numerical results

The initial conditions for saturation are out of equilibrium. The capillary forces set the initially present gas in motion so that the gas gradually spreads downwards, developing the capillary fringe and increasing the dissolution. The system passes through the equilibration stage within the first 39 d (t_{eq}) as the gravity and the capillary pressure are taken

into account. Fig. 5.10 shows the position of the gas front. The fast gas distribution is associated with the negative mass flux through the contact surface between the unsaturated and fully saturated zones, Fig. 5.11.

Meanwhile, convective fluxes start to form over the whole length of the capillary fringe. However, the gas motion is dominant and prevents density fingering. Once the gravity perturbations in the CO₂ rich zone are large enough, the system becomes unstable to the convective fluxes on the transition zone. Its impact can be seen in Fig. 5.11, where starting from the onset time point the mass transfer increases. The onset time, $t_c = 92$ d, can be adjusted assuming $c_0 = 6317$, Eq. (5.3.5). Due to the equilibration stage, the onset time is delayed in comparison with the previous results for the single phase problem (Riaz et al., 2006; Pau et al., 2010). For the relative onset time, $t_c - t_{eq} = 53$ d, $c_0 = 3623$. This value can be overestimated due to the grid dimension. Additional grid refinement would be necessary to define the onset convergence point that was numerically demonstrated by high resolution modeling of single-phase problems (Pau et al., 2010) and also of two-phase problems with transient and stagnant capillary fringe (Elenius et al., 2015). These results correspond to the grid with $h_{x,z} = 1/16$ m. Using the coarser grid provides high underestimation. For example, choosing the grid of 1/16 allow to model 18 fingers during the early convective time, while the grid of 2 m can produce only 1 finger after 5 y.

The CO₂(aq) rich fingers grow down and the mass transfer flux reaches a stabilization stage, approximately at 157 d, Figs. 5.11 and 5.12. During this period, the average mass transfer flux is 9.73×10^{-7} kg/m²/s that is of the same order of magnitude as the results of Pau et al. (2010).

The convective mass transfer is 174 kg of CO₂ during the stabilization period, Fig. 5.12. The density deviation is $\Delta\rho_l = 18.28$ kg/m³. The analytical estimation Eq. (5.3.2) yields 102 kg with $\alpha = 0.2$. The α coefficient adjusted to the mass transfer flux is 0.342. Based on the reported mass fluxes by Pau et al. (2010), the adjusted α is 0.47. The mass flux estimation of Elenius et al. (2014), that also takes into account the capillary fringe, can be fitted to the numerical results with the scaled coefficient 0.055. In Elenius et al. (2015), the mass flux of the models with no transition zone or transient capillary fringe is lower than the estimation of Elenius et al. (2014) with the coefficient 0.03. The adjusted $\alpha = 0.342$ and mass flux can be used to estimate and compare the convective flow at reservoir scale as the deviation of the values is low.

After the onset point, the contact surface between two phases undergoes high perturbations and finally becomes uneven; further, its deviation (maximum and minimum distance) increases with the time. Fig. 5.10 shows that the maximum and minimum gas front vertical position diverges from the outbreak point, 141 d. Therefore, the convective development, Fig. 5.13, leads to a heterogeneous gas front, Fig. 5.14. The uneven gas surface was also exposed in Elenius et al. (2015) by modeling a long term injection in the reservoir scale.

Hence, in addition to the convective motion in the aqueous phase, the instabilities growing on the rough contact surface between two phases imply corresponding perturbations in the gas phase, evolving the gas mixing, Fig. 5.15. It is worth mentioning that the convective mixing in both phases was also observed at the reservoir scale, Sec. 3.3.3.

We believe that the moving uneven transition zone affects the convection both in the aqueous and in the gas phases. The constant gas source also impacts to the process: supplying fresh $\text{CO}_2(\text{g})$ allows for the continued development of new fingers, compare Figs. 5.9 and 5.13. Elenius et al. (2015) have shown that both transient and stagnant capillary transition zones accelerate the onset time and convective mass transfer.

Table 5.2 – Parameters of the 2D problem of the convective mixing.

Initial Conditions		
$\Omega = [0, 10] \times [0, H] \text{ [m}^2\text{]}$		
$\Omega_w = [0, 10] \times [0, H - h_z] \text{ [m}^2\text{]}$		
$\Omega_g = [0, 10] \times [H - h_z, H] \text{ [m}^2\text{]}$		
$h_z = 1/16, 1/8, 1/4, 2 \text{ [m]}$, see Sec. 5.3.1		
$\Omega_w :$	$p_{l0} = p_{atm} + \rho_l g(z_D - z) \text{ [Pa]}$	$S_{g0} = 0$
$\Omega_g :$	$p_{l0} = p_{atm} + \rho_l g(z_D - z) \text{ [Pa]}$	$S_{g0} = 0.4$
Temperature	50 °C	
Boundary Conditions		
$\Gamma_{top} = [(0, H); (10, H)] \text{ [m]}$		
$\Gamma_{top} :$	constant CO ₂ (aq) concentration	
$\partial\Omega :$	no flow	
Matrix Properties		
Porosity ϕ	0.2	
Permeability K	$10^{-12} \text{ [m}^2\text{]}$	
Fluid-Matrix Properties		
Brooks-Corey capillary pressure and relative permeability		
Eq. 2.1.7, Eq. (2.1.9), Eq. (2.1.10): $p_b = 10^5 \text{ Pa}$, $\lambda = 2$, $S_{lr} = 0.15$, $S_{gr} = 0.05$		
Fluid Properties		
Density	ρ_l : T-VTPR	ρ_g : PR78
Viscosity	$\mu_l = 5.49 \times 10^{-5}$	$\mu_g = 2.84 \times 10^{-5} \text{ [Pa} \cdot \text{s]}$
Diffusion coefficient	$D_l = 3.36 \times 10^{-9}$	$D_g = 1.95 \times 10^{-7} \text{ [m}^2\text{/s]}$

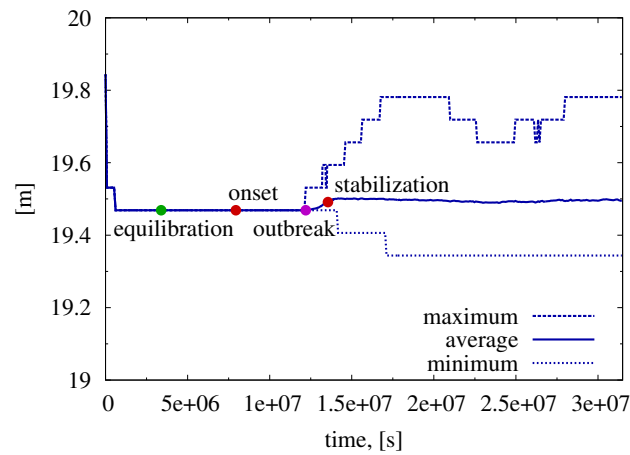


Figure 5.10 – Evolution of the gas front vertical position.

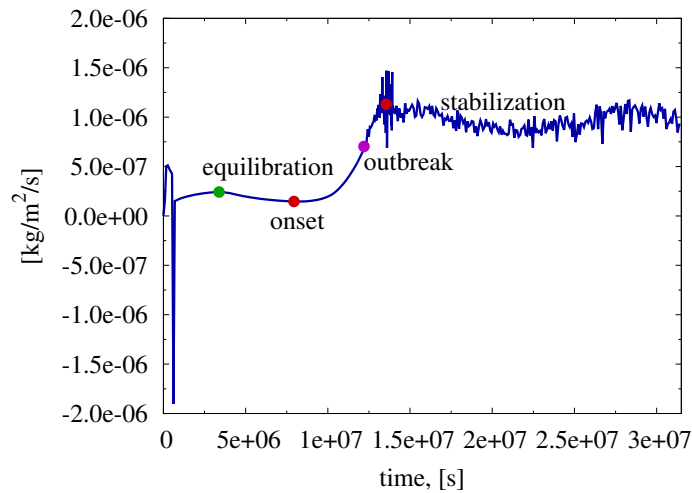


Figure 5.11 – Evolution of the mass flux of CO_2 through the moving contact surface.

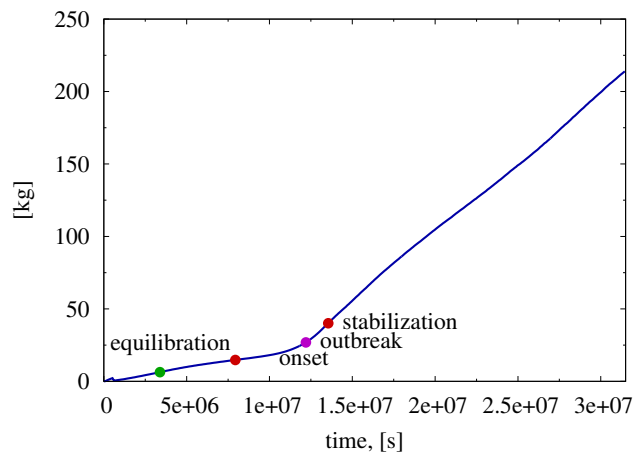


Figure 5.12 – Evolution of the total mass of CO_2 beneath the moving contact surface.

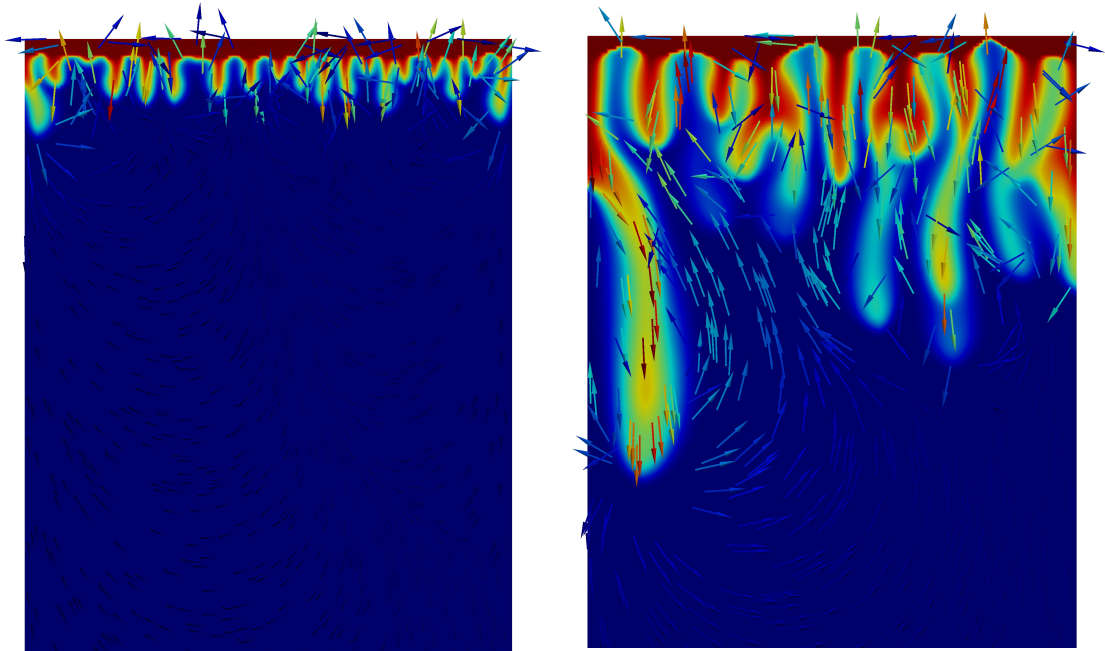


Figure 5.13 – $\text{CO}_2(\text{aq})$ concentration and the velocity field at 120 and 365 d, the length is 10 cm.

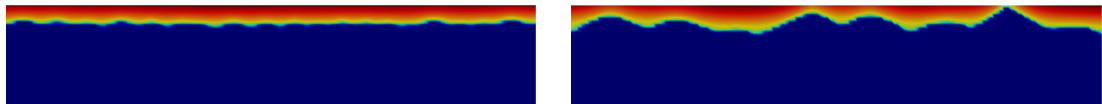


Figure 5.14 – Gas saturation at 120 and 365 d, the length is 10 cm.

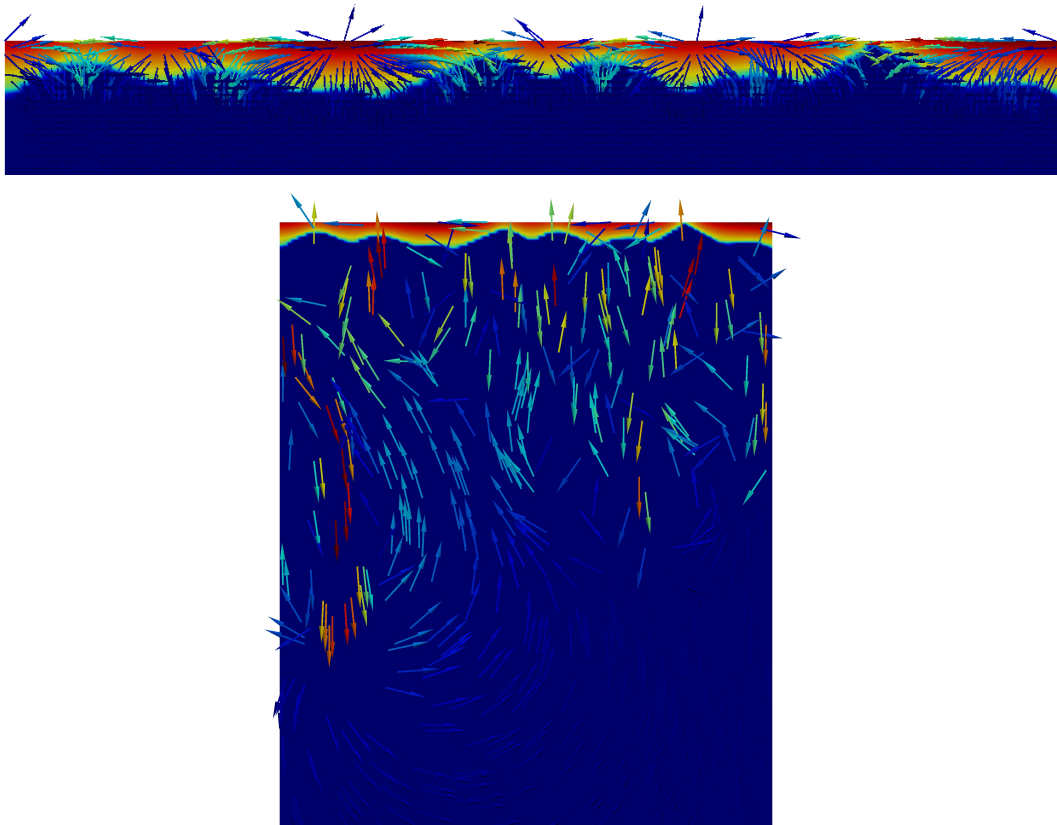


Figure 5.15 – Gas saturation and the velocity field in gas (top) and liquid (bottom) phases at 365 d, the length is 10 cm.

5.4 Modeling CO₂ injection

The modeling of CO₂ injection at the reservoir scale provides an analysis of the gas plume evolution and undergoing processes. However, using the coarse grid results in lower convective fluxes and consequently, in lower dissolution rate. We study the convective underestimation on the radial geometry applying numerical modeling and analytical formulations.

5.4.1 Problem description

The supercritical CO₂ is injected through a well in a vertically confined long aquifer with a height of 100 m at a depth of $z_D = 1100$ m. The injection rate is constant, 10 kg/s; the well radius is 0.3 m; the injection is performed over the whole area of the well during 30 y. Supposing a homogeneous medium, the 2D axisymmetric geometry is chosen. The fluid and matrix parameters are identical to those of the convective mixing problem, Table 5.2. The length of the aquifer is 10 km. Variable cell length from 2 to 1000 m allows to reduce the grid dimension to 33,750 cells.

5.4.2 Numerical results

The injected CO₂(g) rises up and then spreads along the top of the aquifer, Fig 5.16. The maximum expansion can be followed by the radius of this plume at the top of the aquifer, Fig. 5.17. The evolution is proportional to \sqrt{t} , consistent with the axisymmetric geometry. Notice that gas displacement is the dominant mechanism: the dissolution rate is weak, so that the dissolved CO₂ area expansion is limited by the gas displacement.

In the absence of convection, the amount of dissolved CO₂ is proportional to the amount of injected CO₂(g), Fig. 5.18. Indeed, the dissolution occurs only where CO₂(g) is available, and at equilibrium. On the contrary, the convective flux increases CO₂ dissolution by



Figure 5.16 – Gas saturation map in range $[0, \max(S_g)]$: $\max(S_g) = 0.7$ at 120 d (top), $\max(S_g) = 0.73$ at 16 y (center) and 30 y (bottom), $[100, 2500]$ m.

exporting $\text{CO}_2(\text{aq})$ rich water out of the gas current. Consequently, there is a discrepancy between two models after about 120 d, Fig. 5.18. The onset time of the problem is associated to the moment when the current near the well flows downwards. This time point is underestimated due to the grid dimension.

Following these perturbations, the convective fingering occurs at the upper part of the current starting from the region close to the well. The onset time range given in Sec. 5.3.1 does not correspond to the two-phase problem conditions with the moving contact zone.

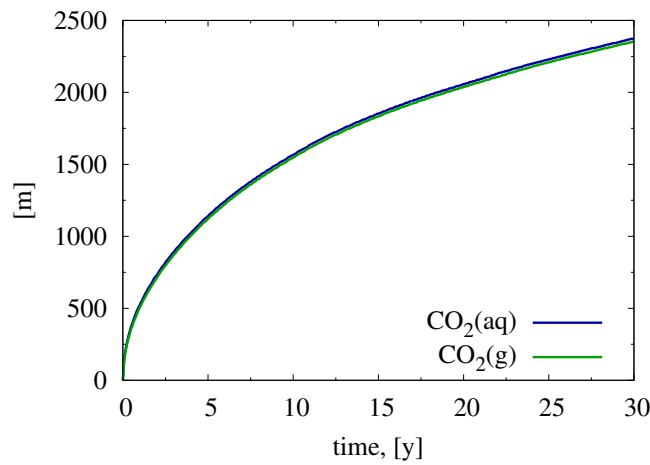


Figure 5.17 – Evolution of the $\text{CO}_2(\text{g})$ and $\text{CO}_2(\text{aq})$ plume radius at the top of the aquifer.

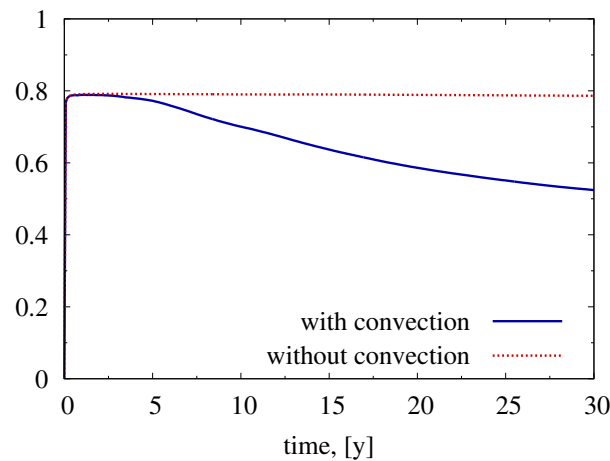


Figure 5.18 – Ratio of the CO_2 quantity in gas phase to the injected $\text{CO}_2(\text{g})$ with and without convection.

The CO₂ rich fingers directed downward provoke the counter water stream. The induced circulation of fresh water enhances the dissolution. In this period, the convection impact is linear with time. Then at about 16 y, we can observe the later convective mixing period, Fig. 5.18, when the liquid under the current contains the dissolved CO₂. At 30 y, 52% of injected CO₂ remains in the gas phase. So, almost half of the injected CO₂ is trapped by dissolution. Without the convective dissolution, the rate of CO₂(g) is nearly constant, 79%.

Figs. 5.19, 5.20 and 5.21 illustrate the CO₂(aq), HCO₃⁻ concentration and pH at 120 d, then 16 y, 30 y corresponding to the different stages of convective mixing. The gas and liquid densities are presented in Fig. 5.22. The gas density gradually grows with increasing hydrostatic pressure. The liquid density deviation is 18.51 kg/m³.

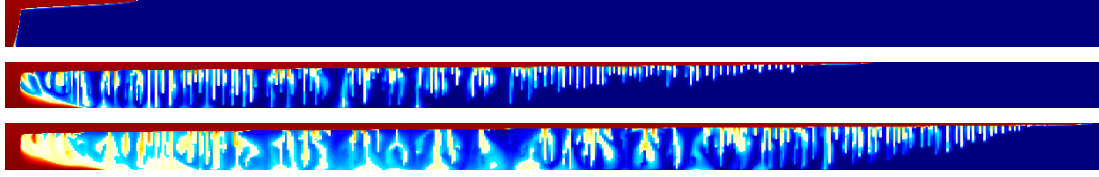


Figure 5.19 – $\text{CO}_2(\text{aq})$ concentration map in range $[0, \max(\text{CO}_2(\text{aq}))]$: $\max(\text{CO}_2(\text{aq})) = 1.296$ molal at 120 d (top), $\max(\text{CO}_2(\text{aq})) = 1.294$ molal at 16 y (center) and 30 y (bottom), $[100, 2500]$ m.

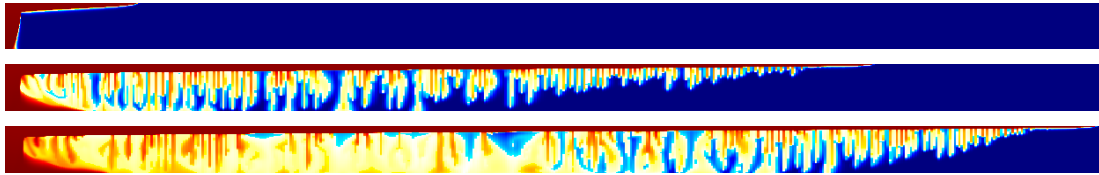


Figure 5.20 – HCO_3^- concentration map at 120 d (top), 16 y (center) and 30 y (bottom), $[100, 2500]$ m. The HCO_3^- scale is $[0, 0.00087]$ molal.

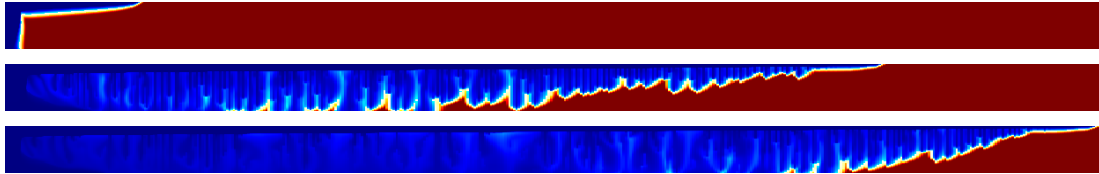


Figure 5.21 – pH map at 120 d (top), 16 y (center) and 30 y (bottom), $[100, 2500]$ m. The pH scale is $[3.07, 6.63]$.

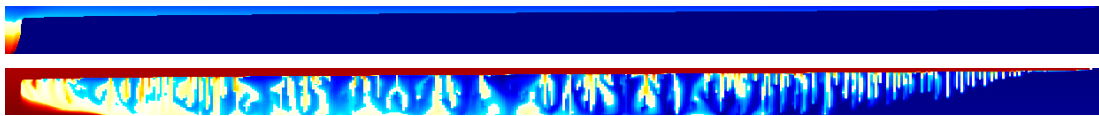


Figure 5.22 – Gas (top) and liquid (bottom) density map at 30 y, $[100, 2500]$ m. The gas and liquid density scales are $[395, 493]$ and $[1022, 1040]$ kg/m^3 , respectively.

5.4.3 Gas current spreading

The numerical results provide the lower convective dissolution at the reservoir scale, so we give the order of this underestimation. The radius of the gas current at the top of the aquifer neglecting dissolution can be expressed by following (Nordbotten et al., 2005):

$$R = \sqrt{\frac{\lambda_g Q t}{\phi \pi \lambda_w H}}, \quad (5.4.1)$$

where λ_g and λ_w are the mobilities of gas and water, Q is the volumetric injection rate. Assuming $\lambda_\alpha = 1/\mu_\alpha$, the analytical estimation Eq. 5.4.1 corresponds well to the numerical results of the plume radius that include the dissolution and diffusion processes, Fig. 5.23. This good agreement entails that neither dissolution nor diffusion has significant impact on the radius of the gas spreading.

Since the CO₂(aq) rich current follows the gas spreading (Fig. 5.17), we imply the convective flux formulation of Neufeld et al. (2010), Eq. (5.3.2), to the radius formula, Eq. (5.4.1) by piecewise time discretization. For the convective impact estimation, we neglect the area near the well and suppose that the shape of current is a disc. Assuming $\alpha = 0.346$, the analytical and numerical results with the convective dissolution perfectly

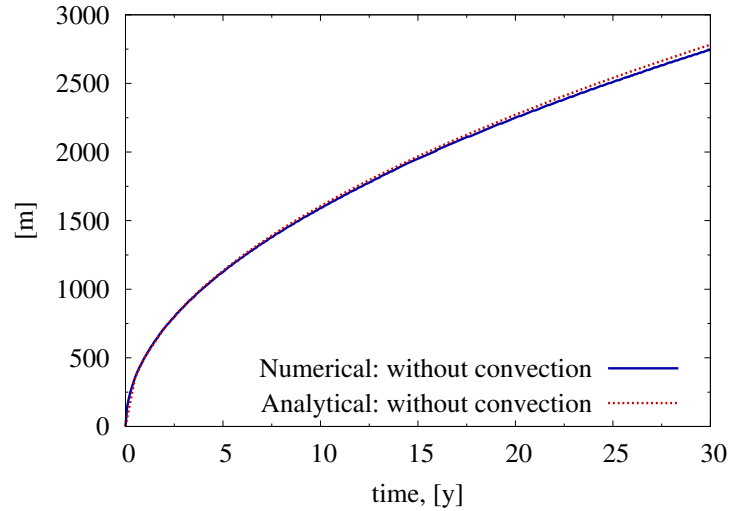


Figure 5.23 – Prediction of current radius at the top of the aquifer without convective dissolution by analytical and numerical methods. The analytical estimation also neglects diffusion and dissolution, Eq. (5.4.1); the numerical results account for diffusion and dissolution.

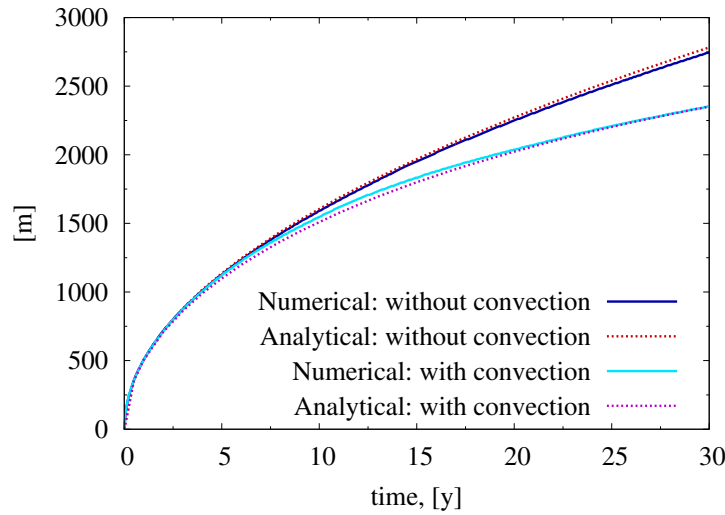


Figure 5.24 – Prediction of current radius at the top of the aquifer with and without convective dissolution by analytical and numerical methods. The analytical solution without convective mixing is given by Eq. (5.4.1); the analytical estimation with convective dissolution is obtained by time piecewise approximation of non-convective analytical solution Eq. (5.4.1) with the convective transfer Eq. (5.3.2).

fit together, Fig. 5.24. This analytical approximation predicts 29% of convective volume, that is 29% of CO_2 is dissolved by the convective mixing. Based on the numerical results, the convective impact is 26%, Fig. 5.18. Given the results in Sec. 5.3.2, it implies a similar deviation of mass transfer flux.

The numerical convective mass transfer is of the same order of magnitude as the estimation of Neufeld et al. but higher by 182%. At the same time, it is smaller than the high resolution numerical results of Pau et al. (2010), about 77% of their value. Moreover, the numerical results of the convective mass transfer are close to those of the higher resolution in Sec. 5.3.2. Pau et al. showed the nearly equivalent mass fluxes on 2D and 3D grid. We thus deduce that the 2D radial modeling can be efficiently used reducing the grid dimension and enhancing the convective mixing in comparison to the 3D modeling.

5.5 CO₂ with impurities and convective mixing

Preliminary observations of the current spread and the impact of impurities:

- From the chromatographic partitioning analysis Sec. 5.2, we conclude that species less soluble than CO₂ promote the advance of the plume; more soluble species reduce the radius. These results have been observed at the field scale, on the pilot in Tongliao, Fig. 5.25.

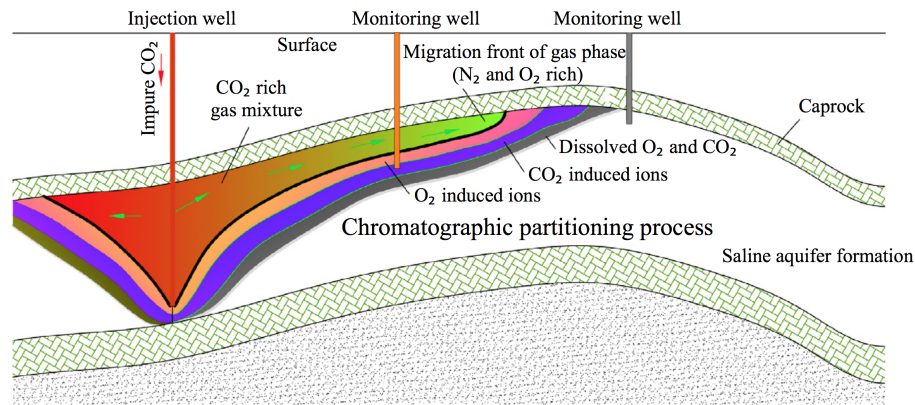


Figure 5.25 – Graphical representation of the pilot in Tongliao, Inner Mongolia, China (Wei et al., 2015).

- The difference between the injected fluid density and that of the aquifer water ($\Delta\rho_{ph} = \rho_{fw} - \rho_{inj}$) influences the distance of the plume spread: the smaller difference implies smaller buoyancy forces, hence the smaller spread radius and vice versa, by Archimedes' principle.
- The difference between the increased density of the aquifer/formation water with the solutes and the initial aquifer water density ($\Delta\rho_l = \rho_{diss} - \rho_{fw}$) is responsible for the density driven convective flow:
 - The higher the density difference, the higher the rate of convective fluxes should be and vice versa, see the experimental and analytical results of Kneafsey and Pruess (2010), MacMinn et al. (2012), Neufeld et al. (2010).
 - The higher the density difference, the smaller the wavelength and the earlier the onset time are expected (Ennis-King et al., 2003; Riaz et al., 2006).

- The higher the convective rate, the faster the aquifer water under the plume becomes saturated by the dissolved gases that decelerates the dissolution rate afterwards and hence extends the plume lifetime if the aquifer height is limited, (Riaz et al., 2006; MacMinn et al., 2012).

The viscosity ratios also affect on the current dynamics. We point out some factors, although the viscosity is supposed to be constant in this study. The spreading increases with $M_{ph} = \mu_{fw}/\mu_{inj}$, where μ_{fw} is the viscosity of the aquifer water, μ_{inj} is the viscosity of the injected gas. The viscosity ratio $M_l = \mu_{fw}/\mu_{diss}$ also affects the current dynamics, μ_{diss} is the viscosity of the aquifer water with the solutes. MacMinn et al. (2012) showed analytically and experimentally that the current extension decreased with M_l lengthening its life-time.

5.5.1 Problem description

Taking into account the chemical and thermodynamic properties of gas impurities, we model the co-injections of SO_2 and Air that have contrary characteristics of solubility, density, chromatographic partitioning effect, Sec. 5.2. Three gas injection scenarios are proposed

- pure CO_2 ;
- 95 mol% of CO_2 with 5 mol% of SO_2 ;
- 95 mol% of CO_2 with 5 mol% of Air (i.e., N_2 and O_2 , 4 and 1 mol% of the total gas composition, respectively).

The mass injection rate is identical, 10 kg/s and performed through the whole height of the aquifer during 10 y. The problem parameters and geometry are identical to those of the pure CO_2 injection model, Sec. 5.4.1. The coarse grid is used to perceive the general dynamics of the system and its sensitivity to the injected gas composition. The cell length is variable, from 4 m to 1 km.

5.5.2 Numerical results

Given the equal mass injection rate and different gas composition, the gas volumetric rate q_{vol} of three scenarios can be written by following

$$q_{vol,SO_2} < q_{vol,CO_2} < q_{vol,Air}, \quad (5.5.1)$$

as their densities are in the inverse order. As a results, the larger gas extension (maximum radius) is observed in the case of Air admixture and the smaller radius in the case of SO₂ admixture. Fig. 5.26 displays the maximum gas radius of the three cases. We stress that the order of plume extensions is the consequence of the gas composition density.

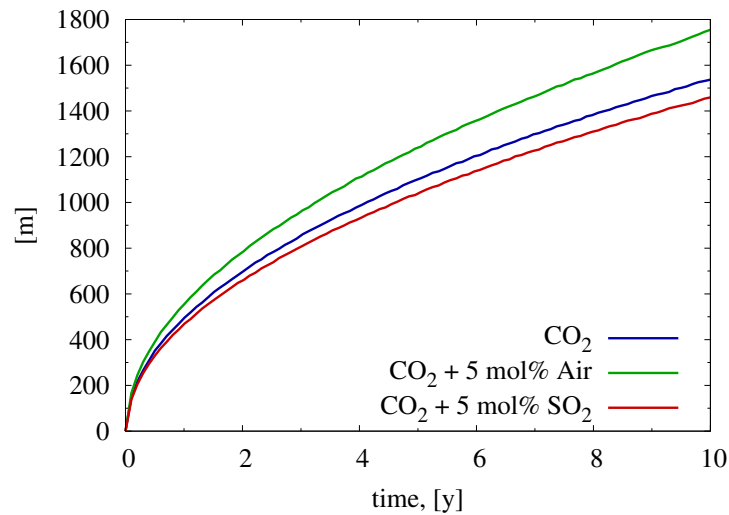


Figure 5.26 – Gas extension at the top of the aquifer (maximum radius of the gas current) for the injection scenarios of pure CO₂, {CO₂ + Air} and {CO₂ + SO₂}.

The gas current density in the case of Air admixture is low in contrast to that of SO₂. In addition, the front of the plume primarily consists of N₂ and O₂ due to the chromatographic partitioning. On the contrary, the co-injection of SO₂ entails the higher gas density and CO₂ rich gas front of the plume, Figs. 5.27 and 5.28.

Since the gas current of the Air case spreads faster than those of pure CO₂ and SO₂ cases, the CO₂ dissolution rate is the highest. Fig. 5.29 illustrates the ratio of CO₂(g) quantity to its total injection mass during 10 y. Similarly to Sec. 5.4.2, the curves decline and converge with time, indicating a growing impact of convection on the dissolution, which is more important in the case of SO₂. Fig. 5.30 displays ratio of CO₂ dissolved with

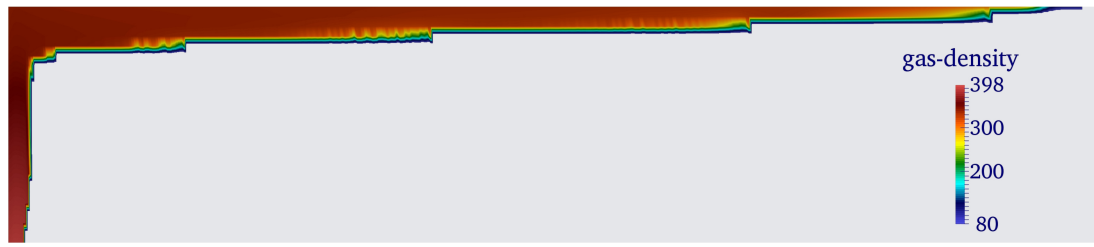


Figure 5.27 – Injection of $\{\text{CO}_2 + \text{Air}\}$: gas density at 10 y, [100,1800] m. (The x-axis is scaled, 4 : 1.)

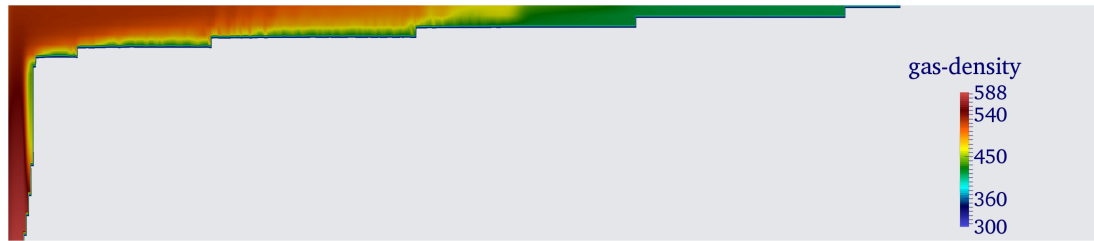


Figure 5.28 – Injection of $\{\text{CO}_2 + \text{SO}_2\}$: gas density at 10 y, [100,1800] m. (The x-axis is scaled, 4 : 1.)

convection to that without convection: the curves thus quantify the impact of convection for each scenario.

The convective processes can be also observed in the gas phase. Fig. 5.31 presents the mole fraction profiles in gas phase at 10 y. The gas convective instabilities of the Air scenario are emphasized by the fact that N_2 and O_2 constitute a thin front in contrast to the advanced CO_2 rich layer in the SO_2 case. As was mentioned in Sec. 5.3.2, this gas behaviour is associated with the convective flow in the liquid phase, Figs. 5.32 and 5.33. The impact of $\text{H}_2\text{O}(\text{g})$ is negligible.

The higher convective mass transfer was expected in the SO_2 scenario (Li and Jiang, 2014; Ziabakhsh-Ganji and Kooi, 2014) as the density difference $\Delta\rho_l$ is almost two times bigger than that of the pure CO_2 and Air scenarios. However, due to the chromatographic partitioning a CO_2 rich front develops. So the convective transfer is similar to that of the pure CO_2 injection. As a result, the curves of convective impact in the pure CO_2 and SO_2 cases are very close, Fig. 5.30.

For the Air scenario, even though the liquid density deviation $\Delta\rho_l$ is smaller than that of the pure CO_2 case, the gas density and chromatographic partitioning of Air scenario are

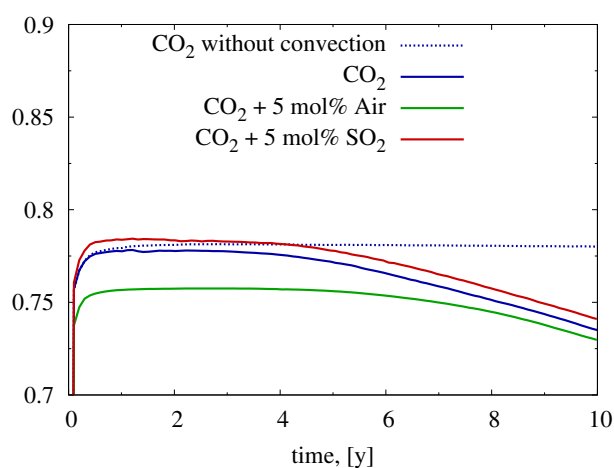


Figure 5.29 – Ratio of the total CO₂(g) quantity to the injected CO₂ for the injection scenarios of pure CO₂, {CO₂ + Air} and {CO₂ + SO₂} during 10 y.

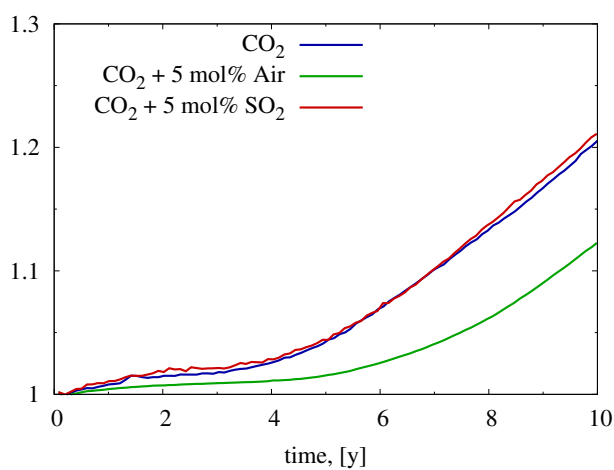


Figure 5.30 – Convective impact for the injection scenarios of pure CO₂, {CO₂ + Air} and {CO₂ + SO₂}: ratio of total dissolved CO₂ with and without convection.

favorable to the gas current advancing, hence to the pure dissolution. Also, it increases the spread area slightly enhancing the convective impact. That explains a low convective impact and high dissolution rate at the same time.

In the light of these results, we conclude that the high interaction between the chromatographic properties of the mixture on the one hand and its convective potential on the other, requires careful modeling two-phase compositional flow and thermodynamics of mixtures. In order to establish further tendency of convective contribution, the injection time should be extended, the grid dimension and refinement is necessary.

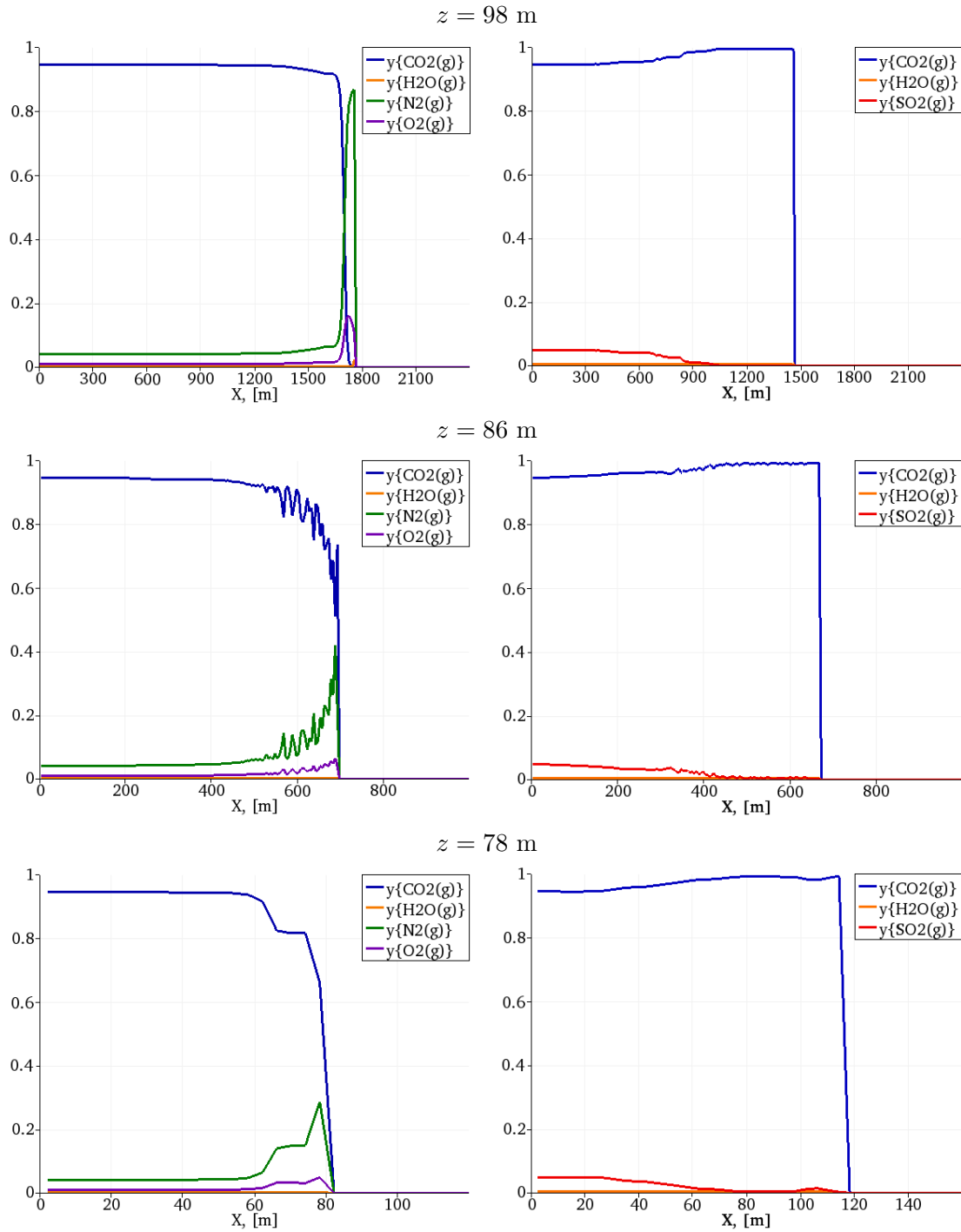


Figure 5.31 – Mole fractions in gas phase at the height 98 (top), 86 (center) and 78 m (bottom) for the problem of $\{CO_2 + Air\}$ (left) and $\{CO_2 + SO_2\}$ (right) injection at 10 y.

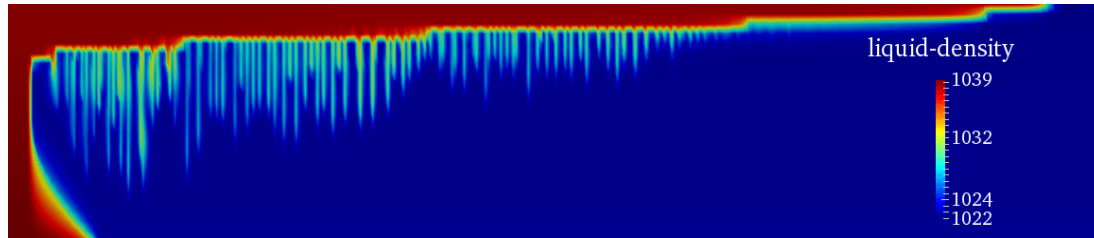


Figure 5.32 – Injection of $\{\text{CO}_2 + \text{Air}\}$: liquid density at 10 y, [100, 1800] m. (The x-axis is scaled, 4 : 1.)

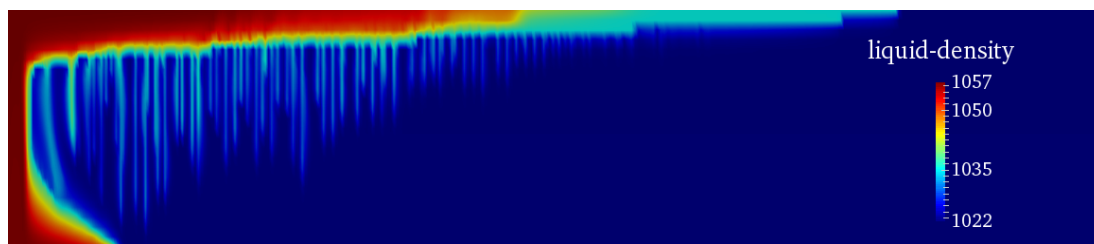


Figure 5.33 – Injection of $\{\text{CO}_2 + \text{SO}_2\}$: liquid density at 10 y, [100, 1800] m. (The x-axis is scaled, 4 : 1.)

Chapter 6

Conclusions and Future Work

Le travail a porté sur les méthodes numériques de couplage du transport réactif et de l'écoulement multicomposant multiphasique compressible. Le dernier est associé à la résolution d'un système non-linéaire dont dimension croît avec le nombre de composants. Pour éviter cette contrainte, une nouvelle approche de couplage par séparation d'opérateurs a été proposée pour faciliter intégration de l'écoulement multiphasique dans le cadre des codes de transport réactif. Puisque la formulation de conservation des phases est employée et couplée au transport compositionnel linéaire, la résolution de l'écoulement peut être séparée ce qui permet de conserver la structure du code de base.

Le module d'écoulement diphasique a été implémenté dans HYTEC ce qui complète ses fonctionnalités de régime saturé et non-saturé (l'équation de Richards). Nous avons opté pour le schéma complètement implicite et jacobien analytique afin de résoudre l'écoulement. Avec l'approche proposée, l'écoulement multicomposant compressible a été couplé au transport réactif. Les équations d'état ont été utilisées pour la modélisation de l'équilibre de phases et la prédiction des propriétés volumétriques dont l'analyse comparative a mis en évidence les avantages de certains modèles en fonction de la composition et des conditions de système.

La vérification du système couplé a reposé sur la comparaison des résultats numériques et analytiques avec ceux obtenus par d'autres codes. Un exercice de benchmarking a été proposé pour la vérification et validation des codes de couplage. Les développements de ce travail ont permis d'effectuer la modélisation de stockage du carbone et d'étudier l'impact de mélange convectif et de présence des impuretés dans le gaz injecté à la dynamique du

système et la dissolution de gaz globale.

Les nouvelles fonctionnalité d'HYTEC ouvrent un large éventail d'applications des géosciences ainsi que de possibilités et questions relatives aux développements futurs du code.

6.1 Conclusions

The present work provides a study of reactive transport coupling with multiphase multi-component compressible flow. A new coupling method was proposed, relying on an operator splitting approach, so that its integration into an existing reactive transport code is facilitated. The method was implemented in the HYTEC code.

Before this work, HYTEC possessed saturated or unsaturated (Richards' equation) flow capabilities, coupled with aqueous and gaseous (diffusive) transport and chemistry. A two-phase flow module was developed, using a fully implicit (with an analytical Jacobian) pressure/saturation algorithm. Then, the operator splitting methodology was implemented, so that multiphase multicomponent compressible flow with complex fluid properties is now coupled with multiphase reactive transport. Applying the formulation of phase conservation allowed to minimize the dimension of flow problem and make its structure composition independent. The coupling method conserved the existing functionality and flexibility of the code. The equations of state were used for the phase equilibria and prediction of volumetric properties, whose comparative analysis highlighted advantages of several models according to the composition and system conditions.

The verification of the code was performed by modeling problems with analytical and literature results; in addition, a benchmark exercise for the multiphase flow and reactive transport modeling community was proposed to verify and validate the coupling methods. The developments of this work were applied for modeling the CO₂ underground storage. The proposed coupling allows to investigate a complex gas injection (CO₂ and impurities) and to reveal new behaviors of gas and liquid dynamics – high interaction between density convective flux and chromatographic partitioning. We investigated the impact of the injected gas composition on the velocity of gas saturation front, density driven fingering and its effect on the overall gas dissolution. This simulation paves the way for future work, including hydrodynamic heterogeneities or pronounced effect of the host-rock mineralogy.

6.2 Future work

The new functionality in HYTEC opens a wide array of applications, as detailed above. It is also obvious that the present work unveils a whole world of questions relative to the future development of the code.

6.2.1 Applications

- Behavior of complex gas for geoscience applications: CO₂ storage but also new forms of storage for the energy sector, like H₂ or high frequency natural gas storage. The behavior of residual air during the saturation stage of nuclear waste facilities is also interesting: although the pressure temperatures are less extreme, O₂ consumption, CO₂ production or consumption over acidic or basic fronts in the storage, H₂ production by the corrosion of steel containers, make for complex gas evolution in the system.
- The impact of several important parameters has already been addressed here: differential solubility, density. The role of capillary pressure should also be investigated in several relevant applications, particularly in heterogeneous media. Capillary pressure discontinuity indeed raises some numerical developments (van Duijn et al., 1995).

6.2.2 Simplifying hypotheses

Along the development of our method and its implementation, several simplifying hypotheses were chosen. An obvious reason is overall limitations of the code: for instance the chemical module CHESS relies on the presence of water, so that a robust algorithm for water disappearance is pointless. Some hypotheses were considered because they simplify the independent modules without effect on the coupling method (e.g., isotropy).

- An isotropic formulation was chosen, consistent with the usual finite volumes Voronoï scheme in HYTEC. A new vertex centered scheme is now available in HYTEC for the saturated mode. Its extension for the two-phase flow is relatively straightforward, although averaging between adjoining vertices should be carefully addressed (particularly for capillary pressure and relative permeability). This implementation would have no impact on the coupling scheme, due to the operator splitting approach.

- Including the feedback of chemistry on the hydrodynamics is still a challenge for the reactive transport community. The impact of mineral volume change on fluid flow can be addressed by a volumetric source term that can be incorporated in the coupling equations of OS structure. Another problem is the impact of the porosity change on the hydrodynamic properties, intrinsic permeability, relative permeability and capillary pressure. No global satisfying method is available so far, apart from specifically designed empirical laws.
- The creation of a gas phase, due to chemical reaction, in an initially saturated zone should also be addressed with care. A comparison between potential equilibrium pressure and actual fluid pressures is a first indicator that can be applied. The communication with the flow module must again be addressed with care. This problem is particularly important in the propagation of acidic plumes in more or less carbonated environments (e.g., in situ recovery), or in non-isothermal modeling.

6.2.3 Extension of the physical problems

- Dual porosity leads to different discretization, homogenized models of flow system, (Masson et al., 2014a; Jurak et al., 2014), and opens the possibility for future researches in coupled multiphase flow and reactive transport in fractured media. The chemical impact in the flow in fracture can be stirring in connection with the RT and variable porosity/permeability due to the geochemical reactions.
- A non-isothermal module is available in HYTEC: for saturated regime only, the heat and flow solvers are sequentially coupled, then the Darcy velocity and the temperature fields are used within the reactive transport coupling (e.g., modification of the solubility and kinetics constants). Non-isothermal multiphase flow raises an issue of coupling method. The next options should be studied: FIM/IMPES flow + explicit/implicit heat. Applying the explicit scheme entails the definition of associated CFL criteria. In terms of perspective coupling with the reactive transport, the chemical impact might be a key factor in choosing explicit or implicit method.
- Coupling free flow and saturated and two-phase flow in porous medium implies introducing Navier-Stokes equations and Beavers-Jeseph interface condition (Rybak

et al., 2015; Brenner et al., 2014).

6.2.4 Numerical methods

- Applying a strong coupling (iterative procedure of flow, transport, chemistry) needs a comparative study with global implicit method in order to establish its capabilities to reproduce the chemical impact on the flow.
- Primary variables of multiphase flow system: once CHESSE can handle the drying out process, the variable switching is unavoidable and it should be generalized.
- The AIM is of great importance for the large scale modeling since it can significantly reduce the nonlinear system. (The IMPES model is applied for decoupled formulation with pressure equation Sec. 2.2.1.) The linear solvers and preconditioners such as CPR (Cao et al., 2005) can be observed for further development. Moreover, in some applications, the coupling with single-phase flow (domain decomposition) also increases computational efficiency. The flexible structure of the proposed coupling method allows different both formulations and techniques.

Nomenclature

Latin Symbols

a	attraction parameter in Peng-Robinson EOS
a_α	dispersivity in phase α , $[m]$
A_α	mass accumulation of phase α
A_s	specific surface area, $[m^2/m^3 \text{ solution}]$ or $[m^2/kg \text{ mineral}]$
b	van der Waals covolume
c_0	onset time constant of convective instability
c_l^k	total liquid mobile concentration of basis species k , $[mol/kg \text{ } w]$
c_s^k	immobile concentration of basis species k , $[mol/kg \text{ } w]$
c_g^m	gas concentration of basis species m , $[mol/m^3]$
C_i	concentration of primary species i in chemical module
d	dissolution parameter of transport model
D_α	molecular diffusion coefficient of phase α , $[m^2/s]$
D_α^e	effective diffusion coefficient of phase α , $[m^2/s]$
e	evaporation parameter of transport model
f_i^α	fugacity of species i in phase α
\mathbf{F}	residual function
\mathbf{g}	gravitational acceleration vector, $[m/s^2]$
\mathbf{J}	Jacobian
k	kinetic constant, $[mol/m^2/s]$
k_{max}^{fl}	maximum number of iterations in flow coupling
$k_{r\alpha}$	relative permeability of phase α
K	intrinsic permeability, $[m^2]$
K_i	K-value/equilibrium ratio

K_j	equilibrium constant of reaction j
K_s	solubility constant of solid phase
K_i^h	Henry's law constant
L	liquid mole fraction
M	molecular weight, $[kg/mol]$
max	maximum over the modeled domain
n_α	quantity of matter in phase α
\mathbf{n}	normal vector
N_c	number of basis species in chemistry module
N_f	number of fluid phases α
N_g	number of gas species
N_{gr}	grid dimension
N_p	number of phases
N_r	number of independent chemical reactions
N_s	number of species
p_α	liquid/gas pressure, $[Pa]$
p_c	capillary pressure, $[Pa]$
p_b	entry pressure of capillary pressure model, $[Pa]$
P	pressure in thermodynamics, $[Pa]$
P_A	attractive term of pressure in cubic equations of state
P_R	repulsive term of pressure in cubic equations of state
q_α	mass source term of phase α , $[kg/s]$
q_g^m	source term of basis species m in gas phase, $[mol/m^3]$
q_l^k	source term of basis species k in liquid phase, $[mol/kg\ w]$
Q	ion activity product
R	gas constant, $[J/K/mol]$
R_α	reaction term of phase α , $[kg/s]$
R_g^m	reaction term of basis species m in gas phase, $[mol/m^3]$
R_l^k	reaction term of basis species k in liquid phase, $[mol/kg\ w]$
Ra	Rayleigh number
S	concentration of species in chemical module

S_j	concentration of species j in chemical module
S_α	saturation of phase α
$S_{\alpha r}$	residual saturation of phase α
Sh	Sherwood number
t_c	onset time of convective instability, [s]
T	Temperature, °C or K
T_i	total concentration in chemistry module
\mathbf{u}_α	Darcy's velocity of phase α
v	molar volume, [m^3/mol]
V	vapor mole fraction
V_α	volume of porous space occupied by phase α , [m^3]
V_{tot}	total volume, [m^3]
X_α^k	mass fraction of basis species k in phase α
y_i	mole fraction of basis species i in gas phase
z_D	depth
Z	compressibility factor

Greek Symbols

$\alpha = \{l, g\}$	liquid/gas phase
α_{ij}	stoichiometric coefficient
β	power parameter of kinetic model
γ_i	activity coefficient of species i
γ_i^∞	activity coefficient of species i at infinite dilution in a solvent k
$\tilde{\gamma}_i$	assymetric activity coefficient
Γ_{ij}	interface between elements Ω_i et Ω_j
Γ_\cdot^D	Dirichlet boundary
Γ_\cdot^N	Neumann boundary
δ	power parameter of kinetic model
Δ	matrix of binary interaction coefficients of PR EOS
ε_g	gas quantity tolerance in reactive transport coupling
ε_{lin}	tolerance of GMRES

ε_{Nf}	residual function tolerance in flow coupling
ε_{qss}	quasi-stationary state tolerance in flow coupling
ε_{rt}	tolerance in reactive transport coupling
ζ	vector of primary variables of flow system
θ	contact angle between the solid surface and the fluid-fluid interface
λ	parameter of Brooks-Corey model
λ_c	wavelength of convective instability, $[m]$
λ_{tot}	total mobility, $[1/(Pa \cdot s)]$
λ_α	mobility of phase α , $[1/(Pa \cdot s)]$
μ	ratio of displaced phase viscosity to the injected one
μ_α	viscosity of phase α , $[Pa \cdot s]$
ρ_α	mass density of phase α , $[kg/m^3]$
ϕ_α	source velocity of phase α , $[m/s]$
τ_α	tortuosity of phase α
ϕ	porosity
φ_i^α	fugacity coefficient of species i in phase α
ψ_α	inflow/outflow over Neumann boundary, $[kg/m^2/s]$
Ω_i	element i
Ω	acentric factor set
\mathfrak{T}_α	transport operator of phase α
$\ \cdot\ $	Eucliden norm
$\ \cdot\ _\infty$	infinity norm

Abbreviations

AIM	adaptive implicit method
CFL	Courant-Friedrichs-Lewy number
CPA-PR	cubic plus association Peng-Robinson equation of state (Hajiw et al., 2015)
CPR	constrained pressure residual (CPR) preconditioner Cao et al. (2005)
CPR	Peng-Robinson equation of state modified by Coquelet et al. (2004)
CSP	compositional space parametrization method (Voskov and Tchelepi, 2009)
DAE	differential algebraic equations based method

DPPS	decoupled phase pressure and saturation formulation
DPS	decoupled pressure and saturation formulation
DSA	direct substitution approach
EOS	equation of state
FIM	fully implicit method
FVM	finite volume method
GIA	global implicit approach
GPR	Peng-Robinson equation of state modified by Gasem et al. (2001)
GPS	global pressure and saturation formulation
HPR	Peng-Robinson equation of state modified by Haghtalab et al. (2011)
IMPEC	implicit pressure/explicit concentration
MMF	multiphase multicomponent flow
MMRF	multiphase multicomponent reactive flow
NVF	natural variables formulation of compositional flow (Coats, 1980)
ODE	ordinary differential equations based method
OS	operator splitting
PPF	pressure pressure formulation of compositional flow (Angelini, 2010)
PPR78	Peng-Robinson equation of state modified by Jaubert and Mutelet (2004)
PR76	Peng-Robinson equation of state (Peng and Robinson, 1976)
PR78	Peng-Robinson equation of state (Robinson and Peng, 1978)
PS	coupled general phase conservation formulation
PSF	pressures, saturations, fugacities formulation of compositional flow (Lauser et al., 2011)
REV	representative elementary volume
RK	Redlich-Kwong equation of state
RT	reactive transport
SIA	sequential iterative approach
SNIA	sequential non-iterative approach
SRK	Soave-Redlich-Kwong equation of state
TPFA	two point flux approximation
TPR	Peng-Robinson equation of state modified by Twu et al. (1995)

T-VTPR	Peng-Robinson equation of state modified by Ahlers and Gmehling (2001)
VLE	vapor-liquid equilibria
VTPR	Peng-Robinson equation of state modified by Ahlers and Gmehling (2001)

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Appendices

Appendix A

Adaptation of Capillary Pressure Function

The van Genuchten capillary pressure model

- The singularity occurs when $\bar{S}_g \rightarrow 0$

$$\lim_{\bar{S}_g \rightarrow 0} p'_c(\bar{S}_g) = \infty.$$

Let us fix \bar{S}_{+0} such that $\bar{S}_{+0} \in (0, \delta)$. Considering that $p_c|_{\bar{S}_g=0} = 0$, a small quadratic spline can be constructed. The user-defined fitting point \bar{S}_{+0} is 0.001 by default, the input parameter is *Sfit0*.

- Despite the restriction of present formulation and

$$\lim_{\bar{S}_g \rightarrow 1} p_c(\bar{S}_g) = \infty,$$

the capillary pressure is redefined at $\bar{S}_g = 1$. By taking S_m such that $S_m < 1 - S_{lr} \leq 1$, the linear extension on $[S_m, 1]$ with a constant derivative $\left(\frac{\partial p_c}{\partial S_g}\right)_{S_g=S_m}$ is added. The fitting point S_m corresponds to input parameter *Sfit1*, 0.001 by default.

Appendix B

Adaptation of Relative Permeability Function

The van Genuchten liquid relative permeability model

- The singularity at $\bar{S}_g \rightarrow 0$ is fixed by redefinition, similar to that of VG capillary pressure at $\bar{S}_g \rightarrow 0$.

$$\lim_{\bar{S}_g \rightarrow 0} k'_{rl}(\bar{S}_g) = \infty$$

Recall $k_{rl} |_{\bar{S}_g=0} = 1$. Once the fitting point S_{gr+0} , $0 \leq S_{gr} < S_{gr+0}$, is chosen, we build the quadratic spline on $[S_{gr}, S_{gr+0}]$. The fitting point S_{gr+0} corresponds to S_{fit0} , 0.001 by default.

The van Genuchten gas relative permeability model

- To avoid the singularity when $\bar{S}_g \rightarrow 0$

$$\lim_{\bar{S}_g \rightarrow 0} k'_{rg}(\bar{S}_g) = \infty$$

the fitting point $S_{fit0} = 1 \times 10^{-16}$ was chosen to reconstruct k_{rg} and the derivative in $[S_{gr}, S_{gr} + S_{fit0}]$.

Modélisation numérique d'écoulement diphasique compressible et transport réactif en milieux poreux — Applications à l'étude de stockage de CO₂ et de réservoir de gaz naturel.

Résumé : Les activités humaines dans la subsurface se développent rapidement (stockage de déchets, nouvelles techniques minières, stockage à haute fréquence de l'énergie), alors que les attentes du public et des autorités s'intensifient. L'évaluation de chaque étape de ces opérations souterraines nécessite des études détaillées de la sûreté et des impacts environnementaux. La modélisation multiphysique permet de comprendre et de prévoir le comportement des systèmes complexes, à différentes échelles de temps et d'espace.

Le but de ce travail est d'intégrer la résolution de l'écoulement diphasique compressible dans le cadre de codes de transport réactif à l'aide d'une méthode de séparation d'opérateurs. Un module multiphasique a été créé dans le code de transport réactif HYTEC. Une nouvelle approche a ensuite été développée pour coupler écoulement multicomposant multiphasique compressible, description de propriétés thermodynamiques des fluides, avec des codes de transport réactif. Une méthode alternative a été proposée pour résoudre séparément le transport multiphasique en utilisant des termes de taux de réaction numériques. Le système couplé a été vérifié et comparé à d'autres formulations de couplage pour la précision et la performance numérique. Un exercice de benchmark basé sur la modélisation d'un réservoir de gaz naturel a été proposé pour examiner les méthodes de couplage de simulateurs multiphasiques réactifs, leurs capacités numériques et caractéristiques.

Une partie de ce travail s'est focalisée sur la modélisation numérique et analytique de captage et stockage du carbone, l'impact de mélange convectif et de présence des impuretés dans le gaz injecté. La méthode de couplage a permis d'étudier des scénarios d'injection de gaz complexes, incluant sans s'y limiter des mélanges de quatre gaz, et de révéler de nouveaux comportements de la dynamique des fluides, gazeux et liquides — interaction entre le flux convectif densitaire et la séparation chromatographique.

Mots clés : Écoulement diphasique compressible, Transport réactif, Couplage par séparation d'opérateurs, HYTEC, Équation d'état, Stockage de CO₂ impur, Mélange convectif.

Numerical simulation of compressible two-phase flow and reactive transport in porous media — Applications to the study of CO₂ storage and natural gas reservoirs.

Abstract: Human activity in the subsurface has been rapidly expanding and diversifying (waste disposal, new mining technologies, high-frequency storage of energy), while the public and regulatory expectations keep growing. The assessment of each step of underground operations requires careful safety and environmental impact evaluations. Multiphysics modeling provides an effective way to understand and predict the behavior of such complex systems at different time and space scales.

This work aims at incorporating a compressible multiphase flow into the conventional reactive transport framework by an operator splitting approach. A multiphase flow module was therefore implemented in the reactive transport software HYTEC using a new approach, developed to fully couple multiphase multicomponent compressible flow and description of the fluid thermodynamic properties with existing reactive transport codes. An alternative method based on the sequential iterative approach was then invented to separately model multiphase transport by means of numerical reaction rate terms. The coupled system was verified and compared with other coupling formulations for accuracy and computational performance. A benchmark exercise of modeling a natural gas reservoir was proposed to investigate the coupling methods of reactive multiphase simulators, as well as their numerical capacity and characteristics.

The final part of this work concerns the numerical and analytical modeling of carbon capture and storage and the impact of convective mixing and gas impurities in the injected stream in particular. The proposed coupling allowed to study complex gas injection scenarios, modeling a system including but not limited to four gases, and to reveal new behaviors of gas and liquid dynamics, specifically the interaction between density convective flux and chromatographic partitioning.

Keywords: Compressible two-phase flow, Reactive transport, Operator splitting based coupling, HYTEC, Equation of state, Impure CO₂ storage, Convective mixing.

